Refine Search

Search Results -

Terms	Documents
L4 AND PESTICIDE	44

US Pre-Grant Publication Full-Text Database US Patents Full-Text Database

US OCR Full-Text Database

EPO Abstracts Database Database:

JPO Abstracts Database

Derwent World Patents Index

IBM Technical Disclosure Bulletins

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Search History

DATE: Tuesday, July 31, 2007 **Purge Queries** Printable Copy Create Case

Set Name side by side	Query	Hit Count	Set Name result set
DB=PGPB	,USPT,USOC,EPAB,JPAB,DWPI,TDBD; PLUR=1	YES; OP=ADJ	
<u>L6</u>	L4 AND PESTICIDE	44	<u>L6</u>
<u>L5</u>	L4 AND PESTCIDE	0	<u>L5</u>
<u>L4</u>	MALONONITRIL\$7.TI.	328	<u>L4</u>
<u>L3</u>	MALONONITRIL\$7	4001	<u>L3</u>
DB=PGPB	; PLUR=YES; OP=ADJ	•	
<u>L2</u>	20060004092	1	<u>L2</u>
<u>L1</u>	2006004092	0	<u>L1</u>

END OF SEARCH HISTORY

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First Hit Clear Generate Collection Fwd Refs Bkwd Refs Generate OACS

Search Results - Record(s) 1 through 10 of 44 returned.

1. Document ID: US 20070117854 A1

L6: Entry 1 of 44

File: PGPB

May 24, 2007

PGPUB-DOCUMENT-NUMBER: 20070117854

PGPUB-FILING-TYPE:

DOCUMENT-IDENTIFIER: US 20070117854 A1

TITLE: Malononitrile compounds and use thereof

PUBLICATION-DATE: May 24, 2007

INVENTOR-INFORMATION:

NAME CITY STATE

Mitsudera; Hiromasa Toyonaka-shi JP

US-CL-CURRENT: 514/383; 548/267.4

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KMC Draw De

2. Document ID: US 20060004092 A1

L6: Entry 2 of 44 File: PGPB

Jan 5, 2006

COUNTRY

PGPUB-DOCUMENT-NUMBER: 20060004092

PGPUB-FILING-TYPE:

DOCUMENT-IDENTIFIER: US 20060004092 A1

TITLE: Malononitrile compound and use thereof pesticides

PUBLICATION-DATE: January 5, 2006

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY

Okada; Satoshi Takarazuka-shi JP
Oohra; Daisuke Toyonaka-shi JP
Otaka; Ken Iwaki-shi JP

US-CL-CURRENT: 514/521; 558/441

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KMC Draw. D

☐ 3. Document ID: US 20050209323 A1

L6: Entry 3 of 44

File: PGPB

Sep 22, 2005

PGPUB-DOCUMENT-NUMBER: 20050209323

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20050209323 A1

TITLE: Malononitrile compounds and their use as pesticides

PUBLICATION-DATE: September 22, 2005

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY

Otaka, Ken Osaka JP
Oohira, Daisuke Osaka JP
Okada, Satoshi Takarazuka-shi JP

US-CL-CURRENT: 514/520; 558/388

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KMAC	Draw, De

4. Document ID: US 20050176784 A1

L6: Entry 4 of 44. File: PGPB Aug 11, 2005

PGPUB-DOCUMENT-NUMBER: 20050176784

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20050176784 A1

TITLE: Malononitrile compounds and their use as pesticides

PUBLICATION-DATE: August 11, 2005

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY

Otaka, Ken Iwaki-shi JP
Oohira, Daisuke Toyonaka-shi JP
Takaoka, Daisuke Toyonaka-shi JP

US-CL-CURRENT: <u>514/357</u>; <u>546/320</u>

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KWIC	Drawi De
							1 1 1 1 1 1 1 1					

5. Document ID: US 20040143007 A1

L6: Entry 5 of 44 File: PGPB Jul 22, 2004

PGPUB-DOCUMENT-NUMBER: 20040143007

PGPUB-FILING-TYPE: new

Record List Display Page 3 of 5

DOCUMENT-IDENTIFIER: US 20040143007 A1

TITLE: Malononitrile compounds and their use as pesticides

PUBLICATION-DATE: July 22, 2004

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY

Otaka, Ken Toyonaka-shi JP
Oohira, Daisuke Minoo-shi JP
Suzuki, Masaya Nishitokyo-shi JP

US-CL-CURRENT: 514/520; 558/409

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KWIC	Drawe D
							,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,					

6. Document ID: US 20040142821 A1

L6: Entry 6 of 44 File: PGPB Jul 22, 2004

PGPUB-DOCUMENT-NUMBER: 20040142821

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20040142821 A1

TITLE: Pesticide composition comprising malononitrile compounds

PUBLICATION-DATE: July 22, 2004

INVENTOR-INFORMATION:

NAME CITY . STATE COUNTRY

Otaka, KenToyonaka-shiJPSuzuki, MasayaNishitokyo-shiJPOohira, DaisukeMinoo-shiJP

US-CL-CURRENT: 504/296; 504/309

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sequences	Attachments	Claims	KWIC	Draw, Dr

7. Document ID: US 20040138065 A1

L6: Entry 7 of 44 File: PGPB Jul 15, 2004

PGPUB-DOCUMENT-NUMBER: 20040138065

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20040138065 A1

TITLE: Malononitrile compounds and their use as pesticides

PUBLICATION-DATE: July 15, 2004

Record List Display

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY

Otaka, Ken Osaka JP
Oohira, Daisuke Osaka JP
Okada, Satoshi Hyogo JP

US-CL-CURRENT: 504/309; 558/388

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KMC Draw. De

8. Document ID: US 7026340 B2

L6: Entry 8 of 44 Fi

File: USPT Apr 11, 2006

US-PAT-NO: 7026340

DOCUMENT-IDENTIFIER: US 7026340 B2

TITLE: Malononitrile compounds and their use as pesticides

PRIOR-PUBLICATION:

DOC-ID

DATE

US 20050176784 A1

August 11, 2005

Full	Title	Citation	Front	Review	Classification	Date	Reference	"我""我看你说话。	production of the same	Claims	KWIC	Draw, De

☐ 9. Document ID: US 7011838 B2

L6: Entry 9 of 44

File: USPT

Mar 14, 2006

US-PAT-NO: 7011838

DOCUMENT-IDENTIFIER: US 7011838 B2

TITLE: Malononitrile compounds and their use as pesticides

PRIOR-PUBLICATION:

DOC-ID

DATE

US 20040138065 A1

July 15, 2004

Full	Title	Citation	Front	Review	Classification	Date	Reference	The whater	1967年 · 1	Claims	KMC	Drawu D
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10. Document ID: US 6353126 B1

L6: Entry 10 of 44

File: USPT

Mar 5, 2002

US-PAT-NO: 6353126

DOCUMENT-IDENTIFIER: US 6353126 B1

TITLE: Process for the production of malononitrile

	Full	Title	Citation	Front	Review	Classification	Date	Reference	MATERIAL	A CHIEBTE	Claims	KMC	Draw, De
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Search Results - Record(s) 11 through 20 of 44 returned.

11. Document ID: US 6297393 B1

L6: Entry 11 of 44

File: USPT

Oct 2, 2001

US-PAT-NO: 6297393

DOCUMENT-IDENTIFIER: US 6297393 B1

TITLE: Process for the preparation of malononitrile

Full Title Citation Front Review Classification Date Reference Claims KMC Draw. Do

☐. 12. Document ID: WO 2006122949 A1

L6: Entry 12 of 44

File: EPAB

Nov 23, 2006

PUB-NO: WO2006122949A1

DOCUMENT-IDENTIFIER: WO 2006122949 A1

TITLE: MALONONITRILES AND THEIR USE AS PESTICIDES

Full Title Citation Front Review Classification Date Reference

☐ 13. Document ID: EP 1710234 A1

L6: Entry 13 of 44

File: EPAB

Oct 11, 2006

PUB-NO: EP001710234A1

DOCUMENT-IDENTIFIER: EP 1710234 A1

TITLE: MALONONITRILE COMPOUND AND USE THEREOF

Full Title Citation Front Review Classification Date Reference

14. Document ID: EP 1704143 A1

L6: Entry 14 of 44

File: EPAB

Sep 27, 2006

PUB-NO: EP001704143A1

DOCUMENT-IDENTIFIER: EP 1704143 A1

TITLE: MALONONITRILE COMPOUND AS PESTICIDES

Full Title Citation Front Review Classification Date Reference

15. Document ID: WO 2005068432 A1

L6: Entry 15 of 44 .

File: EPAB

Jul 28, 2005

PUB-NO: WO2005068432A1

DOCUMENT-IDENTIFIER: WO 2005068432 A1

TITLE: MALONONITRILE COMPOUND AS PESTICIDES

Full Title Citation Front Review Classification Date Reference Reference Citation Claims KMC Draw De

☐ 16. Document ID: EP 1555259 A1

L6: Entry 16 of 44

File: EPAB

Jul 20, 2005

PUB-NO: EP001555259A1

DOCUMENT-IDENTIFIER: EP 1555259 A1

TITLE: Malononitrile compounds as pesticides

Full Title Citation Front Review Classification Date Reference Front Citation Claims KMC Draw. De

17. Document ID: WO 2004020399 A1

L6: Entry 17 of 44

File: EPAB

Mar 11, 2004

PUB-NO: WO2004020399A1

DOCUMENT-IDENTIFIER: WO 2004020399 A1

TITLE: MALONONITRILE COMPOUND AND USE THEREOF AS PESTICIDES

Full Title Citation Front Review Classification Date Reference Company Claims KWC Draw De

18. Document ID: WO 2004006677 A1

L6: Entry 18 of 44

File: EPAB

Jan 22, 2004

PUB-NO: WO2004006677A1

DOCUMENT-IDENTIFIER: WO 2004006677 A1

TITLE: MALONONITRILE COMPOUNDS AND THEIR USE AS PESTICIDES

Full Title Citation Front Review Classification Date Reference

19. Document ID: WO 2089579 A1

L6: Entry 19 of 44

File: EPAB

Nov 14, 2002

PUB-NO: WO002089579A1

DOCUMENT-IDENTIFIER: WO 2089579 A1

TITLE: PESTICIDE COMPOSITION COMPRISING MALONONITRILE COMPOUNDS

Full Title Citation Front Review Classification Date Reference

1 20. Document ID: JP 2007055901 A

L6: Entry 20 of 44

File: DWPI

Mar 8, 2007

DERWENT-ACC-NO: 2007-317065

DERWENT-WEEK: 200731

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 $\label{total compound} \begin{tabular}{ll} \textbf{TITLE: Agent useful for preventing pest on livestock, comprises $\frac{malononitrile}{malononitrile}$ compound and metoxadiazone as active ingredients $\frac{malononitrile}{malononitrile}$ and $\frac{malononitrile}{malononitr$

Full	Title	Citation	Front	Review	Classification	Date	Reference	N. Control	37.27.28	Sec. 2.	1	CI.	aims	KMC	Drag
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Search Results - Record(s) 21 through 30 of 44 returned.

21. Document ID: JP 2007031422 A

L6: Entry 21 of 44

File: DWPI

Feb 8, 2007

DERWENT-ACC-NO: 2007-246448

DERWENT-WEEK: 200725

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TITLE: New <u>malononitrile</u> compound useful for controlling pests, insects, mites and nematodes

Full Title Citation Front Review Classification Date Reference

22. Document ID: JP 2007001934 A

L6: Entry 22 of 44

File: DWPI

Jan 11, 2007

DERWENT-ACC-NO: 2007-211549

DERWENT-WEEK: 200722

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TITLE: Composition for controlling pests such as Blattodea insect-pest and flies, contains <u>malononitrile</u> compound and natural pyrethrin as active ingredients

Full Title Citation Front Review Classification Date Reference

Document ID: JP 2007001935 A

L6: Entry 23 of 44

File: DWPI

Jan 11, 2007

DERWENT-ACC-NO: 2007-165230

DERWENT-WEEK: 200717

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TITLE: Pest controller for controlling harmful pests such as Blattodea insect-pest and flies in agriculture and forestry, contains <u>malononitrile</u> compound and ethofenprox as active ingredients

Full | Title | Citation | Front | Review | Classification | Date | Reference | Company | Claims | KMC | Draw, De

24. Document ID: JP 2007001933 A

L6: Entry 24 of 44

File: DWPI

Jan 11, 2007

DERWENT-ACC-NO: 2007-165229

DERWENT-WEEK: 200717

COPYRIGHT 2007 DERWENT INFORMATION LTD

TITLE: Pest controller for controlling harmful pests such as Blattodea insect-pest and flies in agriculture and forestry, contains malononitrile compound and

bifenthrin as active ingredients

Full Title Citation Front Review Classification Date Reference

25. Document ID: JP 2007001932 A

L6: Entry 25 of 44

File: DWPI

Jan 11, 2007

DERWENT-ACC-NO: 2007-154719

DERWENT-WEEK: 200716

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TITLE: Composition for controlling pests such as Blattodea insect-pest and flies,

contains malononitrile compound and empenthrin as active ingredients

Full Title Citation Front Review Classification Date Reference

26. Document ID: JP 2007001931 A

L6: Entry 26 of 44

File: DWPI

Jan 11, 2007

DERWENT-ACC-NO: 2007-154718

DERWENT-WEEK: 200716

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TITLE: Composition for controlling pests such as Blattodea insect-pest and flies,

contains malononitrile compound and imiprothrin as active ingredients

Full Title Citation Front Review Classification Date Reference Company Communication Claims KMC Draw, De

7. Document ID: JP 2007001930 A

L6: Entry 27 of 44

File: DWPI

Jan 11, 2007

DERWENT-ACC-NO: 2007-154717

DERWENT-WEEK: 200716

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TITLE: Composition for controlling pests such as Blattodea insect-pest and flies, contains <u>malononitrile</u> compound and allethrin or prallethrin as active ingredients

Full Title Citation Front Review Classification Date Reference

28. Document ID: JP 2007001929 A

L6: Entry 28 of 44

File: DWPI

Jan 11, 2007

DERWENT-ACC-NO: 2007-154716

DERWENT-WEEK: 200716

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TITLE: Composition for controlling pests such as flies, ticks and moths, contains malononitrile compound and 2,3,5,6-tetrafluoro benzyl ester compound as active

ingredients

Full	Title	Citation	Front	Review	Classification	Date	Reference	STATE	VANGANIJE AV	Claims	KWIC	Draw, D
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П	29.	Docur	nent I	D: JP 2	2007001928	3 A						
L6:	Enti	y 29 c	of 44			File:	DWPI			Jan 11	, 200	7

DERWENT-ACC-NO: 2007-135315

DERWENT-WEEK: 200714

COPYRIGHT 2007 DERWENT INFORMATION LTD

TITLE: Composition for controlling pests such as Blattodea insect-pest, flies, ticks and moths, contains <u>malononitrile</u> compound and 3-phenoxy benzyl ester compound as active ingredients

Full	Title	Citation	Front	Review	Classification	Date	Reference	Sept of the same	Michigan (S)	Claims	KOMC	Draw, De

7. 30. Document ID: CN 1910156 A, WO 2005068432 A1, JP 2005225875 A, EP 1704143 A1, BR 200506447 A, AU 2005205311 A1

L6: Entry 30 of 44

File: DWPI

Feb 7, 2007

DERWENT-ACC-NO: 2005-571136

DERWENT-WEEK: 200743

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TITLE: New <u>malononitrile</u> compounds used e.g. in pesticidal composition for controlling pest and for controlling parasites living outside of livestock

Full	Title	Citation	Front	Review	Classification	Date	Reference	AP USI	1. A. W.	nar ana	Claims	KMC	Dr
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31. Document ID: CN 1910147 A, WO 2005068423 A1, JP 2005225876 A, EP 1710234 A1, AU 2005205298 A1, BR 200506451 A, US 20070117854 A1, KR 2006127968 A

L6: Entry 31 of 44

File: DWPI

Feb 7, 2007

DERWENT-ACC-NO: 2005-563927

DERWENT-WEEK: 200743

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TITLE: New <u>malononitrile</u> compound for preventing pest such as Hemiptera insectpest, planthoppers, leafhoppers, plant louses, stink bugs, greenhouse whitefly and scale insects

Full Title Citation Front Review Classification Date Reference Claims KMC Draw De Claims AMC Draw De Claims

File: DWPI

DERWENT-ACC-NO: 2005-513905

L6: Entry 32 of 44

DERWENT-WEEK: 200641

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TITLE: New <u>malononitrile</u> derivatives, useful in <u>pesticide</u> composition for controlling pests living external to livestock such as cow or horse

Full Title Citation Front Review Classification Date Reference Claims KWIC Draw, De

33. Document ID: JP 2004099597 A

L6: Entry 33 of 44

File: DWPI

Apr 2, 2004

Jul 20, 2005

DERWENT-ACC-NO: 2004-299646

DERWENT-WEEK: 200428

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TITLE: Pest controller for controlling pest e.g. harmful arthropods, insects, mites and nematodes, and for preventing ectoparasite in animals e.g. cow, pig, sheep and dog, contains <u>malononitrile</u> derivatives as active ingredient

Full Title Citation Front Review Classification Date Reference

☐ 34. Document ID: JP 2004099593 A

L6: Entry 34 of 44

File: DWPI

Apr 2, 2004

DERWENT-ACC-NO: 2004-299644

DERWENT-WEEK: 200428

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TITLE: New <u>malononitrile</u> compounds useful in exterminating pests, arthropods, insects, mites, nematodes and parasites, is new

Full Title Citation Front Review Classification Date Reference Structure Citation Claims KMC Draw. D

T 35. Document ID: ES 2276090 T3, WO 2004006677 A1, JP 2004099592 A, AU 2003281174 A1, EP 1521528 A1, BR 200312638 A, KR 2005019888 A, US 20050176784 A1, CN 1668195 A, IN 200403066 P4, US 7026340 B2, EP 1521528 B1, DE 60309874 E, DE 60309874 T2

L6: Entry 35 of 44

File: DWPI

Jun 16, 2007

DERWENT-ACC-NO: 2004-191068

DERWENT-WEEK: 200742

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TITLE: New <u>malononitrile</u> compounds useful for controlling pests e.g. insect pests, acarine pests and nematode pests in agriculture and forestry

Full Title Citation Front Review Classification Date Reference Company Claims KWIC Draw Do

П 36. Document ID: AU 2002307747 A1, WO 200290321 A1, EP 1392647 A1, BR 200209481 A, US 20040143007 A1

L6: Entry 36 of 44

File: DWPI

Nov 18, 2002

DERWENT-ACC-NO: 2003-093276

DERWENT-WEEK: 200452

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TITLE: New malononitrile compounds useful for controlling pests e.g. insect pests

Full Title Citation Front Review Classification Date Reference

737. Document ID: AU 2002255313 B2, WO 200290320 A2, EP 1385817 A2, BR 200209532 A, HU 200400033 A2, US 20040138065 A1, AU 2002255313 A1, US 7011838 B2, RU 2274638 C2

L6: Entry 37 of 44

File: DWPI

Feb 1, 2007

DERWENT-ACC-NO: 2003-093275

DERWENT-WEEK: 200735

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TITLE: New malononitrile compounds useful for controlling pests e.g. insect pests

Full Title Citation Front Review Classification Date Reference

38. Document ID: IN 200301743 P4, WO 200289579 A1, JP 2003026510 A, JP 2003026511 A, JP 2003026647 A, EP 1385377 A1, KR 2003092137 A, KR 2003092138 A, KR 2004012807 A, BR 200209461 A, US 20040142821 A1, AU 2002307746 A1, CN 1523958 A, CN 1524071 A, TW 223979 B1, US 20050209323 A1, CN 1639114 A

L6: Entry 38 of 44

File: DWPI

Jan 6, 2006

DERWENT-ACC-NO: 2003-093225

DERWENT-WEEK: 200615

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TITLE: Composition useful for controlling pests comprises a malononitrile compound

Full Title Citation Front Review Classification Date Reference

39. Document ID: DE 60012918 T2, US 6297393 B1, EP 1184369 A1, JP 2002145843 A, EP 1184369 B1, DE 60012918 E

L6: Entry 39 of 44

File: DWPI

Sep 8, 2005

DERWENT-ACC-NO: 2002-040115

DERWENT-WEEK: 200559

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TITLE: Preparation of <u>malononitrile</u> from cyanoacetamide useful as building block for pharmaceuticals and <u>pesticides</u>, e.g. thiamine, adenine comprises porous particulate solid substance as absorbent replacing inorganic salt

Full Title Citation Front Review Classification Date Reference

40. Document ID: DE 60005252 E, EP 1065198 A1, CA 2312514 A1, JP 2001039939 A, CN 1296944 A, US 6353126 B1, EP 1065198 B1

L6: Entry 40 of 44

File: DWPI

Oct 23, 2003

DERWENT-ACC-NO: 2001-184252

DERWENT-WEEK: 200377

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TITLE: Preparation of <u>malononitrile</u> by reacting cyanoacetamide with cyanuric chloride in the presence of a catalytic amount of N,N-dimethylformamide in polar solvent

Full Title Citation Front Review Classification Date Reference

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Te	erms	Documents	
L4	AND PESTICIDE	44	

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Search Results - Record(s) 41 through 44 of 44 returned.

T: 41. Document ID: CN 1035874 C, EP 611751 A1, US 5344955 A, AU 9455247 A, BR 9400605 A, CA 2115908 A, CZ 9400355 A3, JP 06256286 A, SK 9400189 A3, ZA 9401114 A, NZ 250918 A, AU 671334 B, EP 611751 B1, DE 69402092 E, ES 2100657 T3, CN 1107465 A, IL 108678 A, RU 2111960 C1, MX 186472 B, SG 49606 A1, SK 280168 B6, CZ 289315 B6, KR 285611 B, CA 2115908 C, JP 3512844 B2

L6: Entry 41 of 44

File: DWPI

Sep 17, 1997

DERWENT-ACC-NO: 1994-265362

DERWENT-WEEK: 200455

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TITLE: Prepn. of sodium salt of 1-amino-1-cyanamido-2,2- di:cyano ethylene - from malononitrile and sodium di:cyanamide in aprotic, dipolar solvent, at elevated temp.

Full Title Citation Front Review Classification Date Reference Citation Claims KWIC Draw De

42. Document ID: EP 508353 A1, DE 59206635 G, CA 2065765 A, JP 05112535 A, US 5216160 A, EP 508353 B1

L6: Entry 42 of 44

File: DWPI

Oct 14, 1992

DERWENT-ACC-NO: 1992-341662

DERWENT-WEEK: 199636

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TITLE: 4-Amino-2-chloro-5-cyano-6-methylthio-pyrimidine prodn. - by reacting malononitrile and carbon dissulphide with strong base, further reacting with methylating agent condensing with cyanamide and cyclising with hydrochloric acid

Full Title Citation Front Review Classification Date Reference

43. Document ID: DE 2601052 A, BE 837583 A, NL 7600379 A, JP 51095029 A, FR 2297838 A, US 4000314 A, ZA 7600227 A, AT 7600244 A, GB 1475974 A, IL 48847 A, CH 604508 A, CA 1061799 A, SU 708979 A

L6: Entry 43 of 44

File: DWPI

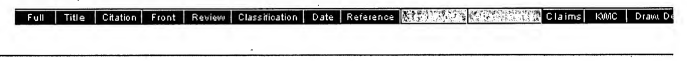
Jul 22, 1976

DERWENT-ACC-NO: 1976-58134X

DERWENT-WEEK: 200300

COPYRIGHT 2007 DERWENT INFORMATION LTD

TITLE: ((N-Benzyl-3-trifluoromethyl-anilino)methylene)malonitrile derivs - prepd by reacting trifluoromethylanilinomethylene_malononitriles with benzyl halides (BE150776)



44. Document ID: US 3726662 A

L6: Entry 44 of 44

File: USOC

Apr 10, 1973

US-PAT-NO: 3726662

DOCUMENT-IDENTIFIER: US 3726662 A

TITLE: HERBICIDAL ANILINOMETHYLENE - MALONONITRILES

DATE-ISSUED: April 10, 1973

INVENTOR-NAME: BAKER J; HOWE R

US-CL-CURRENT: <u>504/312</u>

	Full	Title	Citation	Front	Review	Classification	Date	Reference	Et et la rese	(c) (2) (5)	Claims	KWIC	Drawt D
;	Clear		Genera	ate Co	lection	Print	F	wd Refs	Bkwd	Refs	Gener	ate OA	CŚ
		Te	rms						Docume	nts			
		L4	AND I	PESTI	CIDE						4	14	

Display Format: - Change Format

<u>Previous Page</u> <u>Next Page</u> <u>Go to Doc#</u>

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FILE COVERS 1907 - 31 Jul 2007 VOL 147 ISS 6 FILE LAST UPDATED: 30 Jul 2007 (20070730/ED)

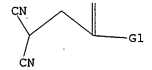
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http://www.cas.org/infopolicy.html

=> d

L1 HAS NO ANSWERS

L1 STR



G1 0, S

Structure attributes must be viewed using STN Express query preparation.

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REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

SAMPLE SEARCH INITIATED 15:33:52 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 63 TO ITERATE

100.0% PROCESSED 63 ITERATIONS

13 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 784 TO 1736
PROJECTED ANSWERS: 44 TO 476

L2 13 SEA SSS SAM L1

=> d 1-10 ibib abs hitstr

L3 ANSWER 1 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2004:203802 CAPLUS

DOCUMENT NUMBER: 140:235428

TITLE: Preparation of malononitrile compound and use thereof

as pesticides

INVENTOR(S): Okada, Satoshi; Oohira, Daisuke; Otaka, Ken PATENT ASSIGNEE(S): Sumitomo Chemical Company, Limited, Japan

SOURCE: PCT Int. Appl., 104 pp.

CODEN: DIVVD2

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	TENT 1	NO.			KIN	D	DATE			APP	LICAT	ION :	NO.		D	ATE	
WO	2004	0203	99		A1	_	2004	0311		WO	2003-	JP10	726		2	0030	826
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BE	, BG,	BR,	BY,	BZ,	CA,	CH,	CN,
		co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	KE,	KG	, KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,
		LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX	, MZ,	NI,	NO,	NZ,	OM,	PG,	PH,
		PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK	, SL,	SY,	TJ,	TM,	TN,	TR,	TT,
											, ZM,			•	·		•
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ	, TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,
											, CH,						
											, NL,						
											, GW,						
AU	2003										2003-					0030	
BR	2003	0139	64		Α		2005	0719		BR	2003-	1396	4		2	0030	826
CN	1678	571			Α		2005	1005		CN	2003-	8204	24		2	0030	826
JP	2004	1431	48		Α		2004	0520		JΡ	2003-	2089	94		2	0030	827
US	2006	0040	92		A1		2006	0105		US	2005-	5227	64		2	0050	201
PRIORIT	Y APP	LN.	INFO	. : `						JΡ	2002-	2503	55		A 2	0020	829
										WO	2003-	JP10	726	1		0030	
OTHER S	OURCE	(S):			MAR	РАТ	140:	23542	2.8								_

OTHER SOURCE(S): MARPAT 140:235428

The present invention relates to a novel malonomitrile commound represented by the formula (I): wherein, R1 represents C1 to C6 alkyl that may be substituted with halogen, C2 to C6 alkenyl that may be substituted with halogen, etc; R2 represents hydrogen atom or C1 to C6 alkyl that may be substituted with halogen; R3 represents hydrogen atom or C1 to C6 alkyl; R4 represents hydrogen atom or C1 to C6 alkyl; R5 represents C1 to C6 alkyl that may be substituted with halogen, C3 to C6 alkenyl that may be substituted with halogen, etc , or R4 and R5 may be combined at their terminal and represent ethylene that may be substituted with C1 to C3 alkyl or trimethylene that may be substituted with C1 to C3 alkyl; and Z1 and Z2, which are the same or different, represent oxygen atom or sulfur atom. Thus, 2-(tert-butoxycarbonylmethyl)-2-allylmalononitrile was prepared by reacting 2-allylmalonitrile with tert-Bu bromoacetate in DMF in the

presence of sodium hydride. The malononitrile compound has an efficient pesticidal activity and can control effectively pests such as insect pests, acarine pests, nematode pests and the like.

IT 666738-88-1P, 2-(tert-Butoxycarbonylmethyl)-2-(1-methylpropyl)malononitrile 666738-93-8P, 2-(tert-Butoxycarbonylmethyl)-2-propylmalononitrile 666738-94-9P, 2-[1-(Ethoxycarbonyl)ethyl]-2-butylmalononitrile 666738-97-2P 666739-14-6P, 2-[(3-Methyl-3-methoxybutoxy)carbonylmethyl]-2-(3,3,3-trifluoropropyl)malononitrile

RL: AGR (Agricultural use); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(production of malononitriles as pesticides)

RN 666738-88-1 CAPLUS

CN Hexanoic acid, 3,3-dicyano-4-methyl-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

RN 666738-93-8 CAPLUS

CN Hexanoic acid, 3,3-dicyano-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

RN 666738-94-9 CAPLUS

CN Heptanoic acid, 3,3-dicyano-2-methyl-, ethyl ester (9CI) (CA INDEX NAME)

RN 666738-97-2 CAPLUS

CN Heptanoic acid, 3,3-dicyano-2-ethyl-, ethyl ester (9CI) (CA INDEX NAME)

RN 666739-14-6 CAPLUS

CN Hexanoic acid, 3,3-dicyano-6,6,6-trifluoro-, 3-methoxy-3-methylbutyl ester

REFERENCE COUNT:

3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 2 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

CORPORATE SOURCE:

2000:141061 CAPLUS

DOCUMENT NUMBER:

132:278722

TITLE:

Spontaneous addition of active methine compounds to

enol ethers and $\dot{\alpha}$, β -unsaturated ketones in

aprotic polar solvent

AUTHOR(S):

Yokozawa, Tsutomu; Oishi, Motoi; Tanaka, Yasukazu Department of Applied Chemistry, Kanagawa University,

Kanagawa-ku Yokohama, 221-8686, Japan

SOURCE:

Journal of Organic Chemistry (2000), 65(6), 1895-1897

CODEN: JOCEAH; ISSN: 0022-3263

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal English

LANGUAGE:
OTHER SOURCE(S):

CASREACT 132:278722.

Addition of (EtO)2CHCH2CXYCH(CN)2 (I, X = Y = cyano; X = cyano, Y = CO2Me; X = Y = CO2Me) to enol ethers and α, β -unsatd. ketones in DMF at room temp is reported. Thus, reacting I (X = Y = cyano) with H2C:CHOEt gave (EtO)2CHCH2C(CN)2CH(OEt)Me in 63% yield. This reaction illustrates that the electron-withdrawing groups at the β -positions of the active methine group having the ones at the α and β positions were strongly affected on the acidity of I.

IT 264142-40-7P 264142-41-8P 264142-43-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (addition of methine compds. to enol ethers and α,β -unsatd. ketones)

RN 264142-40-7 CAPLUS

CN Heptanoic acid, 2,3,3-tricyano-2-(2,2-diethoxyethyl)-6-oxo-, methyl ester (9CI) (CA INDEX NAME)

RN 264142-41-8 CAPLUS

CN Propanedioic acid, (1,1-dicyano-4-oxopentyl)(2,2-diethoxyethyl)-, dimethyl ester (9CI) (CA INDEX NAME)

RN 264142-43-0 CAPLUS

CN Octanoic acid, 2,3,3-tricyano-2-(2,2-diethoxyethyl)-6-oxo-, methyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 3 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1999:142376 CAPLUS

DOCUMENT NUMBER:

130:239567

TITLE:

Diazaspirononanium salt for use as template for

zeolite synthesis

INVENTOR(S):

Kubota, Yoshihiro; Sugi, Yoshihiro

PATENT ASSIGNEE(S): SOURCE:

Showa Denko K. K., Japan Jpn. Kokai Tokkyo Koho, 9 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

1 ~

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
			,	
JP 11060577	Α	19990302	JP 1997-220414	19970815
PRIORITY APPLN. INFO.:		•	JP 1997-220414	19970815
OTHER SOURCE(S):	MARPAT	130:239567	, •	

AB Claimed template is a salt of substituted 2,7-diazaspiro[4,4]nonanium. Hydrothermal synthesis of a zeolite by bringing a silica source and/or an alumina source into contact with the zeolite is also claimed. ZSM-12 zeolites having crystal size of a major axis ≥50 µm are also

claimed.

IT 77415-69-1P

RL: IMF (Industrial manufacture); RCT (Reactant); PREP (Preparation); RACT (Reactant or reagent)

(reaction of; diazaspirononanium salts as templates for manufacture of ZSM-12 zeolites having large crystal size)

RN 77415-69-1 CAPLUS

CN Pentanedioic acid, 3,3-dicyano-, diethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|cccc} \text{O} & \text{CN} & \text{O} \\ \parallel & \parallel & \parallel \\ \text{EtO-C-CH}_2\text{-C-CH}_2\text{-C-OEt} \\ \parallel & \parallel \\ \text{CN} \end{array}$$

L3 ANSWER 4 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1995:900662 CAPLUS

DOCUMENT NUMBER:

124:116317

TITLE:

Lanthanum isopropoxide catalyzed addition of activated

nucleophiles to imines

AUTHOR(S):

LANGUAGE:

Yamamoto, Yoshinori; Fukui, Hiroyuki; Honda, Yoshihiro

CORPORATE SOURCE: SOURCE:

Dept. Chem., Tohoku Univ., Sendai, 980-77, Japan Applied Organometallic Chemistry (1995), 9(5 & 6),

467-71

CODEN: AOCHEX; ISSN: 0268-2605

PUBLISHER: DOCUMENT TYPE:

Wiley Journal English

OTHER SOURCE(S):

CASREACT 124:116317

AB The addition of certain activated nucleophiles to activated imines is catalyzed by lanthanum isopropoxide. As activated nucleophiles,

methylmaloninitrile and Me 2-cyanopropanoate can be utilized. Imines having an electron-withdrawing group either at the carbon or at the nitrogen atom of the C:N double bond can be used: for example

N-toluenesulfonylimines, N-(4-methoxycarbonylphenyl)imines and α -imino esters.

IT 173006-25-2P

RL: SPN (Synthetic preparation); PREP (Preparation)

(lanthanum isopropoxide catalyzed addition of activated nucleophiles to imines)

RN 173006-25-2 CAPLUS

CN Butanoic acid, 3,3-dicyano-2-[(1-phenylethyl)amino]-, 5-methyl-2-(1-methyl-1-phenylethyl)cyclohexyl ester, $[1R-[1\alpha[R^*(R^*)],2\beta,5\alpha]]-$ (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L3 ANSWER 5 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1992:83465 CAPLUS

DOCUMENT NUMBER:

116:83465

TITLE:

The regioselectivity of the ring opening of

1-activated or nonactivated 2-alkoxycarbonyl or 2-cyanoaziridines by carbanions of the dicarbonyl

compounds

AUTHOR(S):

Bouayad, Zoheir; Chanet-Ray, Josette; Ducher, S.;

Vessiere, Roger

CORPORATE SOURCE:

Ec. Natl. Super. Chim. Clermont-Ferrand, Univ. Blaise

Pascal, Aubiere, 63177, Fr.

SOURCE:

Journal of Heterocyclic Chemistry (1991), 28(7),

1757-67

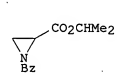
CODEN: JHTCAD; ISSN: 0022-152X

DOCUMENT TYPE: LANGUAGE:

Journal

GT

English



AB Aziridines, e.g. I, reacted with carbanions of dicarbonyl compds., e.g. RO2CCH2CO2R (R = Me, Et, CHMe2), to give ring opened products and/or ring enlarged products, e.g. (RO2C)2CHCH2CH(NHBz)CO2CHMe2, (RO2C)2CHCH(CO2CHMe2)CH2NHBz, and pyrrole II. The regioselectivity depends on several factors. The Ph group on C-3 favors C-3-N bond cleavage, whereas C-2-N bond cleavage is predominant with C-3 substituted or C-2-H aziridines. Cyanoaziridines are predominantly cleaved at C-3-N.

IT 138478-35-0P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

Ι

RN 138478-35-0 CAPLUS

CN Propanoic acid, 2-[(benzoylamino)methyl]-3,3-dicyano-, 1-methylethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \circ \\ || \\ \circ & i\text{-PrO-C} & \text{CN} \\ || & | & | \\ \text{Ph-C-NH-CH}_2\text{-CH-CH-CN} \end{array}$$

L3 ANSWER 6 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1990:497432 CAPLUS

DOCUMENT NUMBER:

113:97432

TITLE:

Quinolone antibacterial agents substituted at the

7-position with spiroamines. Synthesis and

structure-activity relationships

AUTHOR(S):

Culbertson, Townley P.; Sanchez, Joseph P.; Gambino,

Laura; Sesnie, Josephine A.

CORPORATE SOURCE:

Parke-Davis Pharm. Res. Div., Warner-Lambert Co., Annual Co.

Arbor, MI, 48105, USA

SOURCE:

Journal of Medicinal Chemistry (1990), 33(8), 2270-5

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE:

Journal English

LANGUAGE: OTHER SOURCE(S):

CASREACT 113:97432

GI

Fluoroquinolone antibacterials having the 7-position (10-position of AB pyridobenzoxazines) substituted with 2,7-diazaspiro[4.4]nonane,. 1,7-diazaspiro[4.4] nonane, or 2,8-diazaspiro[5.5] undecane (e.g. I (X = CF, CH, N) were prepared and their biol. activities were compared with piperazine and pyrrolidine substituted analogs. Most exhibited potent Gram-pos. and Gram-neg. activity, especially when side chain was N-alkylated. Thus, the quinolinecarboxylic acid II was treated with 2-methyl-2,7-diazaspiro[4.4] nonane to give I (X = CH).

IT 77415-69-1

> RL: RCT (Reactant); RACT (Reactant or reagent) (reductive cyclization of)

RN 77415-69-1 CAPLUS

Pentanedioic acid, 3,3-dicyano-, diethyl ester (9CI) (CA INDEX NAME) CN

ANSWER 7 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1986:496793 CAPLUS

DOCUMENT NUMBER:

105:96793

TITLE:

Zwitterionic tetramethylenes as the common

intermediates in the cycloaddition and polymerization reactions of N-vinylcarbazole with electrophilic tetrasubstituted ethylenes: a new explanation for

charge-transfer initiation

AUTHOR(S):

Gotoh, Tetsuya; Padias, Anne Buyle; Hall, H. K., Jr. Chem. Dep., Univ. Arizona, Tucson, AZ, 85721, USA

SOURCE:

Journal of the American Chemical Society (1986),

108(16), 4920-31

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CORPORATE SOURCE:

CASREACT 105:96793

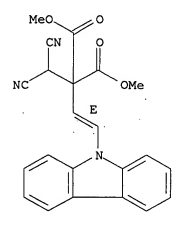
The reactions of N-vinylcarbazole (I) with electrophilic tetrasubstituted ethylenes were examples of reactions whose outcomes are manipulated by changes in concentration, structure, and working procedure to form either small mols. (cyclobutanes, 1-butenes) or poly(vinylcarbazole). Equivalent concns. and evaporative workup (organic chemists' conditions) lead to small mols.; a large excess of I and precipitative workup give polymer. The mechanism involves gauche and trans zwitterionic tetramethylenes as intermediates. The former gives cyclobutane reversibly. The latter gives 1-butenes intramol. or adds monomers to form cyclohexanes or eventually polymer. The organic chemical and polymer chemical are unified on this basis. Extensive stereochem. and kinetic support for these propositions is given. Two other proposed mechanisms for these charge-transfer initiations are excluded.

IT 96735-90-9P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

96735-90-9 CAPLUS RN

CN Propanedioic acid, [2-(9H-carbazol-9-yl)ethenyl](dicyanomethyl)-, dimethyl ester, (E) - (9CI)(CA INDEX NAME)

Double bond geometry as shown.



ANSWER 8 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1985:406747 CAPLUS

DOCUMENT NUMBER: 103:6747

TITLE: Zwitterionic tetramethylene intermediates: a new

interpretation for "charge-transfer" initiation

AUTHOR(S):

Hall, H. K., Jr.; Gotoh, T. Dep. Chem., Univ. Arizona, Tucson, AZ, 85721, USA CORPORATE SOURCE:

SOURCE: Polymer Preprints (American Chemical Society, Division

of Polymer Chemistry) (1985), 26(1), 34-5

医大克二氏管 人名英英格兰克德斯特特

CODEN: ACPPAY; ISSN: 0032-3934

DOCUMENT TYPE: Journal LANGUAGE: English

Investigation of the initiation mechanism in polymerization of N-vinylcarbazole AB [1484-13-5] in the presence of tetracyanoethylene [670-54-2] or di-Me 2,2-dicyanoethylene-1,1-dicarboxylate [82849-49-8] showed that neither the I-cyano compound charge transfer complexes nor the ion-radical pairs formed from them initiated polymerization. The initiating species was the gauche or trans tetramethylene zwitterion formed as an intermediate from the charge-transfer complex. This finding indicated that cyclobutanes initiated vinyl polymerization The mechanism and the kinetics of the zwitterionic initiation were discussed.

IT 96735-90-9

RL: CAT (Catalyst use); USES (Uses)

(catalysts, for vinylcarbazole polymerization)

RN: 96735-90-9 CAPLUS

Propanedioic acid, [2-(9H-carbazol-9-yl)ethenyl](dicyanomethyl)-, dimethyl CN ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

L3 ANSWER 9 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1981:442180 CAPLUS

DOCUMENT NUMBER: 95:42180

TITLE: Absolute configuration of 2,7-diazaspiro[4,4]nonane.

A reassignment

AUTHOR(S): Overberger, C. G.; Wang, David Wei; Hill, Richard K.;

Krow, Grant R.; Ladner, David W.

CORPORATE SOURCE: Macromol. Res. Cent., Univ. Michigan, Ann Arbor, MI,

48109, USA

SOURCE: Journal of Organic Chemistry (1981), 46(13), 2757-64

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 95:42180

GΙ

AB The absolute configuration of the axially dissym. spirane 2,7-diazaspiro[4,4]nonane (I), was elucidated as (R)-(-),(S)-(+) in CHCl3 by synthesis of both enantiomers from the centrodissym. intermediate II; the configuration of (R)-(-)-II was correlated with that of (S)-HO2CCMeEtCH2CO2H through the substituted pyrrolidine III. The configuration thus established for the sulfonamide derivative IV is opposite to that derived earlier (Krow, G. and Hill, R. K., 1968). The source of the original error lies in the preparation of spiroimide V, which is accompanied by almost total racemization when carried out at high temps. A more direct, efficient synthesis of I is described, followed by resolution with dinitrodiphenic acid to give the optically pure enantiomers. Lowe's

rule predicts correctly the absolute configurations of several I derivs. but not that of I itself.

IT 77415-69-1P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and diazaspirononane derivative from)

RN 77415-69-1 CAPLUS

CN Pentanedioic acid, 3,3-dicyano-, diethyl ester (9CI) (CA INDEX NAME)

L3 ANSWER 10 OF 10 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1978:50444 CAPLUS

DOCUMENT NUMBER:

88:50444

TITLE:

The chemistry of 2-oxopropanedinitrile (carbonyl

cyanide); XIX. The ene synthesis using

2-oxopropanedinitrile and 1,3-dicarbonyl compounds

AUTHOR(S):

Kociolek, K.; Leplawy, M. T.

CORPORATE SOURCE:

Inst. Org. Chem., Tech. Univ. Lodz, Lodz, Pol.

SOURCE:

Synthesis (1977), (11), 778-80

DOCUMENT MUDE

CODEN: SYNTBF; ISSN: 0039-7881

DOCUMENT TYPE:

Journal English

LANGUAGE:

Eligitali

OTHER SOURCE(S):

CASREACT 88:50444

AB Reaction of CO(CN)2 with RCOCH2COR1 (I; R = R1 = Ph, 2,4,6-Cl3C6H2, Me; R = Me, F3C, R1 = Ph) in ether at 0° was complete in 1 h and gave RCOCH(COR1)C(CN)2OH (II; R and R1 as before) in 100% yield. Reaction of CO(CN)2 with I (R = R1 = OEt) at room temperature required 20 days and gave II in 43-66% yield.

IT 65305-78-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, with aniline)

RN 65305-78-4 CAPLUS

CN Propanedioic acid, (dicyanohydroxymethyl)-, diethyl ester (9CI) (CA INDEX NAME)

=> s 11 full

REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SEARCH INITIATED 15:35:27 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 1125 TO ITERATE

100.0% PROCESSED 1125 ITERATIONS

SEARCH TIME: 00.00.01

L4 208 SEA SSS FUL L1

L5 78 L4

=> s 15 amd py<2002 MISSING OPERATOR L5 AMD The search profile that was entered contains terms or nested terms that are not separated by a logical operator.

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L6 68 L5 AND PY<2002

.=> d 1-68 ibib abs hitstr

L6 ANSWER 1 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:597989 CAPLUS

DOCUMENT NUMBER: 135:166840

TITLE: Preparation of pyrimidine compounds as modulators of

208 ANSWERS

chemokine receptor activity

INVENTOR(S): Bonnert, Roger; Cage, Peter; Hunt, Fraser; Walters,

Lain; Willis, Paul

PATENT ASSIGNEE(S): Astrazeneca Ab, Swed.

SOURCE: PCT Int. Appl., 52 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA'	rent 1	NO.			KIN	D -	DATÉ			APPL:		ION I			D.	ATE	
WO	2001	0589	02		A1		2001	0816							2	0010	207 <
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
		CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,
		HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,
		LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,
		SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,
		YU,	ZA,	ZW ·				٠.	7. *								
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,
										IT,							
		ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG	•	·
GB	2359	078			Α		2001	0815		GB 2	000-	3019		-	2	0000	211 <
	1265																
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
										AL,				•	·	·	·
JP	2003	5221	89		T		2003	0722	·	JP 2	001-	5580	51		2	0010	207
· US	2003	0405	23		A 1		2003	0227		US 2	002-	2035	84		2	0020	809
บร	6958	344			B2		2005	1025									
US	2005	2340	77		A1		2005	1020		US 2	005-	3668	2		2	0050	114
PRIORIT										GB 2							

OTHER SOURCE(S):

MARPAT 135:166840

GT

$$NR^{2}R^{3}$$
 $NR^{2}R^{3}$
 $NR^{$

AB The title compds. [I; A = II, III (X = NH, CR18R19; Y = N, CR18; R18, R19 = H, alkyl, Ph); R1 = (un)substituted cycloalkyl, alkyl, alkenyl, etc.; R2, R3 = H, cycloalkyl, alkyl, etc.; NR2R3 = (un)substituted 3-8 membered ring optionally containing one or more atoms selected from O, S, NH, etc.], useful in treating an inflammatory disease such as psoriasis and COPD, were prepared E.g., a multi-step synthesis of the 6H-pyrrolo[2,3-d]pyrimidin-6-one IV was given. The compds. I were found to have IC50 of < 10 μM against CXCR2 receptor binding.

IT 224637-77-8

RL: RCT (Reactant); RACT (Reactant or reagent)
 (preparation of pyrimidine compds. as modulators of chemokine receptor
 activity)

RN 224637-77-8 CAPLUS

CN Propanoic acid, 3,3-dicyano-, ethyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT:

6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 2 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2001:347933 CAPLUS

DOCUMENT NUMBER:

135:122094

TITLE:

Cyclopropanation of benzylidenemalononitrile with dialkoxycarbenes and free radical rearrangement of the

cyclopropanes

AUTHOR(S):

Merkley, Nadine; Venneri, Paul C.; Warkentin, John

CORPORATE SOURCE:

Department of Chemistry, McMaster University,

Hamilton, ON, L8S 4M1, Can.

SOURCE:

Canadian Journal of Chemistry (2001), 79(3),

312-318

CODEN: CJCHAG; ISSN: 0008-4042

PUBLISHER:

National Research Council of Canada

DOCUMENT TYPE: LANGUAGE:

Journal English

OTHER SOURCE(S):

CASREACT 135:122094

Thermolysis of 2-cinnamyloxy-2-methoxy-5,5-dimethyl- $\Delta 3$ -1,3,4oxadiazoline (1a) and the analogous 2-benzyloxy-2-methoxy compound (1b) at 110°C, in benzene containing benzylidenemalononitrile, afforded products of apparent regiospecific addition of methoxycarbonyl and cinnamyl (or benzyl) radicals to the double bond. When the thermolysis of la was run with added TEMPO, methoxycarbonyl and cinnamyl radicals were captured. Thermolysis of the 2,2-dibenzyloxy analog (1c) in the presence of benzylidenemalononitrile gave an adduct that is formally the product of addition of benzyloxycarbonyl and benzyl radicals to the double bond. In this case, a radical addition mechanism could be ruled out, because the rate constant for decarboxylation of benzyloxycarbonyl radicals is very large. A mechanism that fits all of the results is predominant cyclopropanation of benzylidenemalononitrile by the dialkoxycarbenes derived from the oxadiazolines, in competition with fragmentation of the carbenes to radical pairs. The cyclopropanes so formed then undergo homolytic ring-opening to the appropriate diradicals. Subsequent β -scission of the diradicals to afford radical pairs, and coupling of those pairs, gives the final products. Thus, both carbene and radical chemical are involved in the overall processes.

IT 351207-62-0P 351207-63-1P 351207-65-3P

351207-66-4P 351207-67-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (cyclopropanation of benzylidenemalononitrile with dialkoxycarbenes and free radical rearrangement of the cyclopropanes)

RN 351207-62-0 CAPLUS

CN Benzeneacetic acid, α -(1,1-dicyano-4-phenyl-3-butenyl)-, methyl ester (9CI) (CA INDEX NAME)

RN 351207-63-1 CAPLUS

CN Benzenebutanoic acid, β , β -dicyano- γ -ethenyl- α -phenyl-, methyl ester (9CI) (CA INDEX NAME)

RN 351207-65-3 CAPLUS

CN Benzenebutanoic acid, β , β -dicyano- α -phenyl-, methyl ester (9CI) (CA INDEX NAME)

RN 351207-66-4 CAPLUS

CN Benzenebutanoic acid, β , β -dicyano- α -phenyl-, phenylmethyl ester (9CI) (CA INDEX NAME)

RN351207-67-5 CAPLUS

CN Benzenebutanoic acid, β , β -dicyano- α -ethoxy- α -methyl-, methyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 3 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:310730 CAPLUS

DOCUMENT NUMBER: 133:104844

New aspects of knoevenagel condensation and michael TITLE:

addition reactions on alkaline carbonates

Aramendia, Maria A.; Borau, Victoriano; Jimenez, AUTHOR(S):

Cesar; Marinas, Jose M.; Romero, Francisco J.

CORPORATE SOURCE: Department of Organic Chemistry, Faculty of Sciences,

Cordoba University, Cordoba, E-14004, Spain

SOURCE: Chemistry Letters (2000), (5), 574-575

CODEN: CMLTAG; ISSN: 0366-7022

PUBLISHER: Chemical Society of Japan

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 133:104844

AB . The Knoevenagel condensation of malononitrile with benzaldehyde on K2CO3. Rb2CO3 and Cs2CO3 gave the condensation product benzylidenemalononitrile but the reaction proceeded to the hydrogenated product benzylmalononitrile. Also, the Michael addition of malononitrile to certain double bonds occurs in the presence of K2CO3.

IT 82584-86-9P

RL: SPN (Synthetic preparation); PREP (Preparation)

(potassium carbonate catalyzed Michael addition reactions of malononitrile with alkenes)

RN 82584-86-9 CAPLUS

CN Butanedioic acid, (dicyanomethyl) -, diethyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT:

12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 4 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:199325 CAPLUS

DOCUMENT NUMBER: 132:237096

TITLE:

SOURCE:

Preparation of 1H-pyrrolo-[1,2-b][1,2,4]triazole

INVENTOR(S):

Morita, Kensuke

PATENT ASSIGNEE(S):

Fuji Photo Film Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 15 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent Japanese

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2000086662 PRIORITY APPLN. INFO.: OTHER SOURCE(S): GI	A CASREA	20000328 ACT 132:23709	JP 1998-265064 JP 1998-265064 96; MARPAT 132:237096	19980918 < 19980918

$$\mathbb{R}^{2} \xrightarrow{\mathbb{N}} \mathbb{N} \mathbb{R}^{1}$$

$$R^2$$
 N
 N
 N
 N
 N
 N
 N

Title compds. I (R1-R3 = H, substituent) are prepared from triazoles II (R1-R3 = H, rsubstituent) via III (R1- R3 = H, substituent) is a substituent 3-(Tert-butylphenyl)-5-[[4-methyl-2,6-di-tertbutylhexyloxycarbonyl]bromomethyl]-1H-1,2,4-triazole was reacted with malononitrile in dimethylacetamide in the presence of NaOMe/MeOH under reflux for 30 min and reacted in the presence of CuCl in PhMe-hexane mixture under reflux for 3 h to give 91% III (R1 = 4-tert-butylphenyl; R2 = cyano, R3 = 4-methyl-2, 6-di-tert-butylhexyloxycarbonyl), which was reacted with isoamyl nitrite in iso-Pr alc. at 50° for 10 h to give 40% I (R1-R3 = same as above).

IT 259266-71-2

RN

RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of pyrrolotriazoles from cyanoethyltriazoles) 259266-71-2 CAPLUS

III

CN lH-1,2,4-Triazole-3-acetic acid, α -(dicyanomethyl)-5-(4-methyl-3-nitrophenyl)-, 2,6-bis(1,1-dimethylethyl)-4-methylcyclohexyl ester (9CI) (CA INDEX NAME)

Me
$$\stackrel{NO_2}{\underset{N-N}{|}}$$
 $\stackrel{CN}{\underset{|}{\underset{t-Bu}{|}}}$ $\stackrel{CN}{\underset{t-Bu}{|}}$ $\stackrel{T-Bu}{\underset{me}{|}}$

IT 259266-70-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of pyrrolotriazoles from cyanoethyltriazoles)

RN 259266-70-1 CAPLUS

CN $1H-1,2,4-Triazole-3-acetic acid, \alpha-(dicyanomethyl)-5-[4-(1,1-dimethylethyl)phenyl]-, 2,6-bis(1,1-dimethylethyl)-4-methylcyclohexylester (9CI) (CA INDEX NAME)$

L6 ANSWER 5 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2000:198039 CAPLUS

DOCUMENT NUMBER:

132:238369

TITLE:

1H-Pyrrolo[1,2-b][1,2,4]triazole derivatives and their

manufacture

INVENTOR(S):

Morita, Kensuke

PATENT ASSIGNEE(S):

Fuji Photo Film Co., Ltd., Japan

Jpn. Kokai Tokkyo Koho, 14 pp. CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

Japanese

FAMILY ACC. NUM. COUNT:

1

PATENT INFORMATION:

PATENT NO.	** KIND	DATE	APPLICATION NO.	DATE * * * * * * * * * * * * * * * * * * *
JP 2000086661 PRIORITY APPLN. INFO.: OTHER SOURCE(S): GI	A MARPAT	20000328	JP 1998-265059 JP 1998-265059	19980918 < 19980918

NC
$$\mathbb{R}^2$$
 \mathbb{R}^3 \mathbb{N} \mathbb{R}^3 \mathbb{R}^3

$$R^2$$
 R^3
 N
 N
 N
 R^1
 N

AB The derivs. III and IV, useful for photog. couplers, physiol. active substances, etc., are manufactured from triazole derivs. I via intermediates II (R1-3 = H, substituent; A = non-metal atomic group to form azole ring with N; X, Y = non-metal atom to form 5-membered ring with CONCO). Thus, I (R1 = p-tert-BuC6H4; R2 = CN; R3 = 2,6-di-tert-butyl-4-methylcyclohexyloxycarbonyl) was treated with CuCl to give 91% II, 9.42 mmol of which was treated with 11.3 mmol acetonylacetone in benzene in the presence of AcOH to give 8.20 mmol III (azole ring = 3,4-dimethyl-1-pyrrolyl).

IT 259266-70-1P

RL: IMF (Industrial manufacture); RCT (Reactant); PREP (Preparation); RACT (Reactant or reagent)

(manufacture of pyrrolotriazole derivs.)

RN 259266-70-1 CAPLUS

CN 1H-1,2,4-Triazole-3-acetic acid, α-(dicyanomethyl)-5-[4-(1,1-dimethylethyl)phenyl]-, 2,6-bis(1,1-dimethylethyl)-4-methylcyclohexyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 6 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2000:141061 CAPLUS

DOCUMENT NUMBER:

CORPORATE SOURCE:

132:278722

TITLE:

Spontaneous addition of active methine compounds to

enol ethers and α, β -unsaturated ketones in

aprotic polar solvent

AUTHOR(S):

Yokozawa, Tsutomu; Oishi, Motoi; Tanaka, Yasukazu Department of Applied Chemistry, Kanagawa University,

Kanagawa-ku Yokohama, 221-8686, Japan

SOURCE:

Journal of Organic Chemistry (2000), 65(6),

1895-1897

CODEN: JOCEAH; ISSN: 0022-3263

PUBLISHER:

American Chemical Society

DOCUMENT TYPE: LANGUAGE:

Journal English

OTHER SOURCE(S):

CASREACT 132:278722

AB Addition of (EtO)2CHCH2CXYCH(CN)2 (I, X = Y = cyano; X = cyano, Y = CO2Me; X = Y = CO2Me) to enol ethers and α , β -unsatd. ketones in DMF at room temp is reported. Thus, reacting I (X = Y = cyano) with H2C:CHOEt gave (EtO)2CHCH2C(CN)2CH(OEt)Me in 63% yield. This reaction illustrates that the electron-withdrawing groups at the β -positions of the active methine group having the ones at the α and β positions were strongly affected on the acidity of I.

IT 184092-93-1 189348-52-5

RL: RCT (Reactant); RACT (Reactant or reagent) (addition of methine compds. to enol ethers and α,β -unsatd. ketones)

RN 184092-93-1 CAPLUS

CN Butanoic acid, 2-cyano-2-(dicyanomethyl)-4,4-diethoxy-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|cccc} \text{CN} & \text{CN} & \text{OEt} \\ & & & & \\ \text{NC-CH-C-CH}_2 - \text{CH-OEt} \\ & & \\ & & \text{C-OMe} \\ & & \\ & & & \\ & & & \\ \end{array}$$

RN 189348-52-5 CAPLUS

CN Propanedioic acid, (dicyanomethyl)(2,2-diethoxyethyl)-, dimethyl ester (9CI) (CA INDEX NAME)

IT 264142-31-6P 264142-33-8P 264142-35-0P

264142-37-2P 264142-39-4P 264142-40-7P

1264142-41-8P 264142-43-0P 264142-45-2P

RL: SPN (Synthetic preparation); PREP (Preparation) (addition of methine compds. to enol ethers and α,β -unsatd. ketones)

RN 264142-31-6 CAPLUS

CN Pentanoic acid, 2,3,3-tricyano-2-(2,2-diethoxyethyl)-4-ethoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 264142-33-8 CAPLUS

CN Pentanoic acid, 4-butoxy-2,3,3-tricyano-2-(2,2-diethoxyethyl)-, methyl ester (9CI) (CA INDEX NAME)

RN 264142-35-0 CAPLUS

CN Pentanoic acid, 4-(2-chloroethoxy)-2,3,3-tricyano-2-(2,2-diethoxyethyl)-, methyl ester (9CI) (CA INDEX NAME)

RN 264142-37-2 CAPLUS

CN 2-Furanpropanoic acid, α, β, β -tricyano- α -(2,2-diethoxyethyl)tetrahydro-, methyl ester (9CI) (CA INDEX NAME)

RN 264142-39-4 CAPLUS

CN 2H-Pyran-2-propanoic acid, α,β,β -tricyano- α -(2,2-diethoxyethyl)tetrahydro-, methyl ester (9CI) (CA INDEX NAME)

RN 264142-40-7 CAPLUS

CN Heptanoic acid, 2,3,3-tricyano-2-(2,2-diethoxyethyl)-6-oxo-, methyl ester (9CI) (CA INDEX NAME)

RN 264142-41-8 CAPLUS

CN Propanedioic acid, (1,1-dicyano-4-oxopentyl)(2,2-diethoxyethyl)-, dimethyl ester (9CI) (CA INDEX NAME)

RN 264142-43-0 CAPLUS

CN Octanoic acid, 2,3,3-tricyano-2-(2,2-diethoxyethyl)-6-oxo-, methyl ester (9CI) (CA INDEX NAME)

RN 264142-45-2 CAPLUS

CN Benzenehexanoic acid, α, β, β -tricyano- α -(2,2-diethoxyethyl)- ϵ -oxo-, methyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT:

25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 7 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2000:139178 CAPLUS

DOCUMENT NUMBER:

132:180579

TITLE:

Preparation of 1H-pyrrolo[1,2-b][1,2,4]triazol-5-

ylamines

INVENTOR(S):

Morita, Kensuke

PATENT ASSIGNEE(S):

Fuji Photo Film Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 15 pp.

SOURCE:

GI

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2000063382	Α	20000229	JP 1998-232925	19980819 <
PRIORITY APPLN. INFO.:			JP 1998-232925	19980819
OTHER SOURCE(S):	CASREA	ACT 132:1805	79; MARPAT 132:180579	•

- AB Title compds. I (R1-R3 = H, substituent), useful as intermediates for physiol. active substances, photog. couplers, dyes, etc., are prepared from triazoles II (X = CHR3CHR2CN; R1-R3 = same as I). II (R1 = C6H4Bu-t-p, X = CHBrCO2Q) was treated with malononitrile and MeONa in DMF-MeOH under ice-cooling for 30 min and heated in the presence of CuCl in PhMe-hexane under reflux for 3 h to give 91% I (R1 = C6H4Bu-t-p, R2 = cyano, R3 = CO2Q).
- IT 259266-70-1P 259266-71-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of pyrrolotriazolylamines as intermediates for physiol. active substances, dyes, and photog. couplers)

RN 259266-70-1 CAPLUS

CN 1H-1,2,4-Triazole-3-acetic acid, α-(dicyanomethyl)-5-[4-(1,1-dimethylethyl)phenyl]-, 2,6-bis(1,1-dimethylethyl)-4-methylcyclohexylester (9CI) (CA INDEX NAME)

RN 259266-71-2 CAPLUS

CN $1H-1,2,4-Triazole-3-acetic acid, \alpha-(dicyanomethyl)-5-(4-methyl-3-nitrophenyl)-, 2,6-bis(1,1-dimethylethyl)-4-methylcyclohexyl ester (9CI) (CA INDEX NAME)$

L6 ANSWER 8 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1999:380253 CAPLUS

DOCUMENT NUMBER:

131:170252

TITLE:

Tandem dimerization and double annulation of 3,3,4,4-tetracyanobutanal acetal. Synthesis of a

bicyclic 2-aminopyridine derivative

AUTHOR(S):

Yokozawa, Tsutomu; Nishikata, Akira; Kimura, Takamasa;

Shimizu, Kazuki; Takehana, Tomoyuki

CORPORATE SOURCE:

Department of Applied Chemistry, Kanagawa University,

Yokohama, 221-8686, Japan

SOURCE:

Tetrahedron Letters (1999), 40(25),

4707-4710

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER:

Elsevier Science Ltd.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 131:170252

GI

AB 3,3,4,4-Tetracyanobutanal acetal (I), which is easily obtained from tetracyanoethylene, Et vinyl ether, and ethanol, yielded 2-aminopyridine derivative II fused with cyclopentane in one pot in the presence of pyridine.

On the basis of several expts., the proposed mechanism involves the Michael reaction of I with the diene generated by the elimination of hydrogen cyanide and ethanol from I, followed by double intramol. nucleophilic addns. to the cyano groups.

IT 184092-93-1

RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of bicyclic aminopyridine by tandem dimerization-cyclization of tetracyanobutanal acetal)

RN 184092-93-1 CAPLUS

CN Butanoic acid, 2-cyano-2-(dicyanomethyl)-4,4-diethoxy-, methyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 9 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1

1999:142792 CAPLUS

DOCUMENT NUMBER:

130:360507

TITLE:

An N2S2 Bifunctional Chelator for Technetium-99m and Rhenium: Complexation, Conjugation, and Epimerization

to a Single Isomer

AUTHOR(S):

SOURCE:

Luyt, Leonard G.; Jenkins, Hilary A.; Hunter, Duncan

Η.

CORPORATE SOURCE:

Department of Chemistry, University of Western

Ontario, London, ON, N6A 5B7, Can. Bioconjugate Chemistry (1999), 10(3),

470-479

CODEN: BCCHES; ISSN: 1043-1802

PUBLISHER:

CODEN. DCC1123, 133N. 1043-100

DOCUMENT TYPE:

American Chemical Society

LANGUAGE:

Journal English

A bifunctional chelator HO2CCH(CH2NHCOCH2SH)2 6 (H3L) was prepared bearing an N2S2 core for binding Re or Tc and a carboxylic acid group for conjugation to amino groups of biomols. Complexation of 6 with Re(V) resulted in two kinetic isomers, anti-and syn-[ReO(HL)]- 7, being formed in approx. equal amts. Epimerization with 0.5M NaOH yields a single isomer anti-7, as determined by NMR spectroscopy and single-crystal x-ray anal. [99mTcO(HL)] - was prepared at the tracer level by reaction of the ligand with 99mTcO4-, SnCl2 and Na gluconate giving a mixture of two isomers, but showing a preference for the anti-isomer. Chelation in the presence of 1^{-1} M NaOH results in anti-8 being formed as the sole product. bifunctional ability of the ligand was explored by amide formation with $(S)-\alpha$ -phenethylamine, either by direct DCC coupling or through the RO2CCH(CH2NHCOCH2STr)2 9 (R = succinimidyl) intermediate. The deprotected bioconjugate PhCHMeNHOCCH(CH2NHCOCH2SH)2 11 (H2L1) was complexed with Re, yielding similar amts. of two isomeric Re complexes, anti- and syn-12, which were isolated and characterized by NMR spectroscopy. Treatment of the kinetic mixture of anti- and syn-[ReOL1]- 12 with 1 M NaOH resulted in quant. conversion to a single Re complex anti-12. With 99mTc in 0.1M NaOAc, bioconjugate 11 yielded anti- and sym-[99mTcOL1]- 13 in a 2:1 ratio, resp. In contrast, complexation in the presence of 1 M NaOH gave only one 99mTc complex, assigned the structure anti-13.

IT 224637-77-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reactant for preparation of rhenium and technetium bis(thioacetamidomethyl)propionate and bis(thioacetamidomethyl)propanam ide complexes)

RN 224637-77-8 CAPLUS

CN Propanoic acid, 3,3-dicyano-, ethyl ester (9CI) (CA INDEX NAME)

O CN | | | | EtO-C-CH₂-CH-CN

REFERENCE COUNT: 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 10 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1999:142376 CAPLUS

DOCUMENT NUMBER:

130:239567

TITLE:

Diazaspirononanium salt for use as template for

zeolite synthesis

INVENTOR(S):

Kubota, Yoshihiro; Sugi, Yoshihiro

PATENT ASSIGNEE(S):

Showa Denko K. K., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 9 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE			
JP 11060577 PRIORITY APPLN. INFO.:	Α	19990302	JP 1997-220414 JP 1997-220414	19970815 < 19970815			
OTHER SOURCE(S):	MARPAT	130:239567	•				

AB Claimed template is a salt of substituted 2,7-diazaspiro[4,4]nonanium. Hydrothermal synthesis of a zeolite by bringing a silica source and/or an alumina source into contact with the zeolite is also claimed. ZSM-12 zeolites having crystal size of a major axis $\geq\!50~\mu m$ are also claimed.

IT 77415-69-1P

RL: IMF (Industrial manufacture); RCT (Reactant); PREP (Preparation); RACT (Reactant or reagent)

(reaction of; diazaspirononanium salts as templates for manufacture of ZSM-12 zeolites having large crystal size)

RN 77415-69-1 CAPLUS

CN Pentanedioic acid, 3,3-dicyano-, diethyl ester (9CI) (CA INDEX NAME)

المرازية أأسأع المدار فالرادات ويريعونيهم والمصهر الإنتصاف الاراطية

L6 ANSWER 11 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1999:118566 CAPLUS

DOCUMENT NUMBER:

130:237421

TITLE:

Stereoselective synthesis of alkenylated malonic

diamide using masked acyl cyanide

AUTHOR(S):

Nemoto, Hisao; Ibaragi, Touru; Bando, Masahiko; Kido,

Masaru; Shibuya, Masayuki

CORPORATE SOURCE:

Faculty of Pharmaceutical Sciences, the University of

Tokushima, Tokushima, 770-8505, Japan

SOURCE:

Tetrahedron Letters (1999), 40(7), 1319-1322

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER:

Elsevier Science Ltd.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 130:237421

GI

AB A highly stereoselective synthesis of an alkenylated malonic diamide (I) starting from a γ , δ -epoxy α , β -unsatd. carboxamide was accomplished using masked acyl cyanide (protected hydroxymalonitrile) via palladium-catalyzed regio- and stereoselective carbon-carbon bond

formation.

ΙT 221219-72-3P 221219-73-4P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(stereoselective synthesis of alkenylated malonic diamide using masked acyl cyanide)

RN 221219-72-3 CAPLUS

3-Hexenoic acid, 2-[dicyano[[(1,1-dimethylethyl)dimethylsilyl]oxy]methyl]-CN 5-hydroxy-, 4-nitrophenyl ester, (2R, 3E, 5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry. Double bond geometry as shown.

221219-73-4 CAPLUS RN

CN 3-Hexenoic acid, 2-[dicyano[[(1,1-dimethylethyl)dimethylsilyl]oxy]methyl]-5-hydroxy-, methyl ester, (2R,3E,5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry. Double bond geometry as shown.

REFERENCE COUNT:

THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 12 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1998:402031 CAPLUS

DOCUMENT NUMBER:

129:122635

TITLE:

The reaction of phenacylmalononitrile with hydrazines:

synthesis of new pyridazinones and

pyrazolo[1,5-a]pyrimidines

AUTHOR(S):

Elnagdi, Mohamed Hilmy; El-Ghamry, Ibrahim; Kandeel, Ezz; Abdel Rahman, A. H.; Al-Naggar, Abdul Aziz; Amer,

Samir; Riad, Mohamed

CORPORATE SOURCE:

Department of Chemistry, Faculty of Science, University of Kuwait, Safat, 13060, Kuwait Gazzetta Chimica Italiana (1997), 127(12),

SOURCE:

azzetta Chimica italiana (1997 01-704

791-794

CODEN: GCITA9; ISSN: 0016-5603

PUBLISHER:

Societa Chimica Italiana Journal

DOCUMENT TYPE: LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 129:122635

AB The reaction of phenacylmalononitrile with hydrazine hydrate affords a mixture of 3,5-diamino-4-phenacylpyrazole (I), and 6-oxo-3-phenyl-1,4,5,6-tetraydropyridazine-5-carbonitrile. The reaction of I with a variety of reagents, that enabled the synthesis of some new pyrazolo[1,5-a]pyrimidine derivs., is described.

IT 210347-41-4P

RL: SPN (Synthetic preparation); PREP (Preparation)

(synthesis of pyridazinones and pyrazolo[1,5-a]pyrimidines)

RN 210347-41-4 CAPLUS

CN Pentanedioic acid, 3,3-dicyano-, bis(1,1-dimethylethyl) ester (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 13 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1998:76233 CAPLUS

DOCUMENT NUMBER:

128:177233

TITLE:

Malononitrile derivatives and herbicides containing

them

INVENTOR(S):

Hosokawa, Akemi; Ikeda, Osamu

PATENT ASSIGNEE(S):

Mitsubishi Chemical Industries Ltd., Japan

Jpn. Kokai Tokkyo Koho, 18 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

KIND DATE APPLICATION NO. DATE

JP 10029966

PATENT NO.

19980203

PRIORITY APPLN. INFO.:

Α

JP 1996-187796 JP 1996-187796 19960717 <--19960717

OTHER SOURCE(S):

MARPAT 128:177233

$$Q =$$
 $E =$
 $E =$

$$Q^{2} = \underbrace{\begin{array}{c} z_{1} \\ CH_{2} \\ E \end{array}}_{E} x_{n}$$

AB The derivs. are represented by R1R2C(CN)2 [I; R1 = H, C1-6 alkyl, C4-7 cycloalkyl, C2-6 alkenyl, C2-6 alkynyl, C1-6 haloalkyl, C2-6 haloalkenyl, C2-6 haloalkynyl, C2-8 alkoxyalkyl, C3-6 alkoxycarbonylalkyl, C2-4 cyanoalkyl, C1-6 hydroxyalkyl, C2-7 alkylamido, C7-9 aralkyl, C8-12 arylcarbonylalkyl, (un)substituted Ph, (un)substituted pyridyl, (un) substituted thiazolyl, CR3R4A; A = (un) substituted Ph, pyridyl, thiazolyl; R3-4 = H, Me; R2 = Q, Q1, Q2; E = CH, N; X = halo, C1-4 alkyl, C1-3 haloalkyl, NO2, C1-8 haloalkoxy, (un) substituted benzyloxy, pyridyloxy; n = 0-2; Z1-2 = OH, halo, C1-4 alkylsulfonyloxy, (un) substituted phenylsulfonyloxy]. The herbicides contain I as active ingredients. I (R1 = H, R2 = Q, E = CH, X = 3-Me, 5-Me) showed 91-100% herbicidal activity against Echinochloa oryzicola, Monochoria vaginalis, and Scirpus juncoides.

IT 203127-60-0P 203127-94-0P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of malononitrile derivs. as herbicides)

RN 203127-60-0 CAPLUS

CN Benzenepentanoic acid, 3-chloro- β , β -dicyano- δ -methylene-, ethyl ester (9CI) (CA INDEX NAME)

RN 203127-94-0 CAPLUS

CN Oxiranebutanoic acid, 2-(3-chlorophenyl)- β , β -dicyano-, ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
CN & O \\
CH_2 - C - CH_2 - C - OEt \\
CN & CN
\end{array}$$

ANSWER 14 OF 68 1.6 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1997:731010 CAPLUS

DOCUMENT NUMBER:

127:346753

TITLE:

Synthesis of terpolymers by spontaneous

copolymerization of the cyclobutane adducts of electron-acceptor olefins and vinyl ether with

2-oxazolines

AUTHOR(S):

Yokozawa, Tsutomu; Tagami, Masato; Takehana, Tomoyuki;

Suzuki, Tadashi

CORPORATE SOURCE:

Dep. Appl. Chem., Kanagawa Univ., Yokohama, 221, Japan

SOURCE: Tetrahedron (1997), 53(45), 15603-15616

CODEN: TETRAB; ISSN: 0040-4020

PUBLISHER:

Elsevier Journal English

DOCUMENT TYPE: LANGUAGE:

AΒ Spontaneous copolymns. of the cyclobutane adducts of strong donors olefins and strong acceptor olefins, 1,1,2,2-tetracyano-3-ethoxycyclobutene (I) and di-Me 2,2-dicyano-3-ethoxycyclobutane-1,1-dicarboxylate (II), with 2-oxazolines are described. In the reaction of II with 2-methyloxazoline (III), the alternating copolymer of II and III, the 1:1:1 periodic terpolymer of di-Me 1,1-dicyanoethylene-2,2-dicarboxylate, vinyl ether, and III, was obtained. Cyclobutane I also reacted with III to yield copolymer rich in I.

IT 198274-09-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and characterization of)

RN198274-09-8 CAPLUS

CN Propanedioic acid, [2-[acetyl[2-(acetyloxy)ethyl]amino]-2ethoxyethyl] (dicyanomethyl) -, dimethyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 15 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

37

ACCESSION NUMBER:

1997:513581 CAPLUS

DOCUMENT NUMBER:

127:184884

TITLE: ·

Multinuclear cluster complexes as diagnostic imaging

contrast agents

INVENTOR(S):

Droege, Michael; Yu, Shi-Bao; Sanderson, William;

Bacon, Edward; Delecki, Daniel; Earley, William; Ye,

Naidong

PATENT ASSIGNEE(S):

Nycomed Salutar, Inc., USA

SOURCE:

PCT Int. Appl., 51 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

PTND

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA:	LENT	NO.			KIN.	D .	DATE				LICAT					ATE		
	9726 9726				A2 A3		1997 1997				1997-					9970	123	<
	W:	AL, DK, LK, RO, KE,	AM, EE, LR, RU, LS,	AT, ES, LS, SD, MW,	AU, FI, LT, SE, SD,	AZ, GB, LU, SG, SZ,	BA, GE, LV, SI, UG,	BB, HU, MD, SK, AT,	BG, IL, MG, TJ, BE,	IS MK TM CH	, BY, , JP, , MN, , TR, , DE, , CF,	KE, MW, TT, DK,	KG, MX, UA, ES,	KP, NO, UG, FI,	KR, NZ, US, FR,	KZ, PL, UZ, GB,	LC, PT, VN GR,	
AU EP	2241 9714 8761 8761	MR, 190 523 61	NE,	SN,	TD, A1 A	TG	1997 1997	0731 0820 1111		CA AU	1997- 1997- 1997-	2241 1452	190 3		1: 1:	9970; 9970; 9970;	123 123	<
CN HU BR	R: 1208 9901 9707 2000 9803	DE, 353 488 300 5158 371	DK,	ES,	FR, A A2 A	GB,	IT, 1999 1999 1999 2000	SE, 0217 0830 1228 1128	•	CN HU BR JP NO GB	1997- 1999- 1997- 1997- 1998- 1996-	1488 7300 5266 3371 1340	68		19 19 19	9970: 9970: 9970: 9980: 9960:	123 123 123 722 123	< <

OTHER SOURCE(S): MARPAT 127:184884

AB Diagnostic imaging contrast media are claimed comprising a physiol. tolerable image contrast-enhancing complex, said complex comprising a pair of interconjugated multinuclear clusters, together with at least one pharmaceutical carrier or excipient. Included, for example, are multinuclear cluster complexes (M3)2L3 containing three metal atoms and L is a ligand. Clusters (M3)2L3 include M3 = M3SaOb where a = 1-4, b = 0-3 and a + b = 4, e.g., M3 = W3SO3. Ligands L include various

polyaminocarboxylates and derivs. represented by general formula (R2)2N[(CHR4)mNR1]n(CHR4)mN(R2)2, e.g., N'-serinol-, N'-methyl-, and N'-benzyl-N,N,N'',N''-diethylenetriaminetetraacetic acids, various N'-(polyhydroxyalkyl)-N'-methyldiethylenetriaminetetraacetic acids, 2-carboxymethylpropylenediaminetetraacetic acid, etc., for which prepns. are given of these and other example ligands. Preparation of cluster compds., e.g., Na4[(W3SO3)2(EGTA)3] (EGTA = ethyleneglycol bis(2-aminoethyl ether)-N,N,N',N'-tetraacetate), from [W3SO3(H2O)9]Cl4 and the appropriate polyaminocarboxylic acid ligand, are described. The claimed preparation of [W3SO3(H2O)9]Cl4 comprises reaction of W(CO)6 and Na2S, followed by acidification of the product with at least 6 N HCl, and purification A charged contrast medium complex may be post-complexed with, e.g., cholamine hydrochloride or N-methyl-N, N-bis (hydroxyethyl) ethylenediamine, to give a preferred neutral derivative Pharmaceutically acceptable forms of the diagnostic imaging contrast media comprising said cluster complexes and dosages for the x-ray contrast media are briefly discussed.

IT 194083-97-1P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(for preparation of multinuclear tungsten aminocarboxylate cluster complexes as diagnostic imaging contrast agents)

RN 194083-97-1 CAPLUS

CN Propanoic acid, 3,3-dicyano- (9CI) (CA INDEX NAME)

CN $NC-CH-CH_2-CO_2H$

ANSWER 16 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1997:414890 CAPLUS

DOCUMENT NUMBER:

127:144690

TITLE:

Metabolism and disposition of the antifolate LY231514

in mice and dogs

AUTHOR(S):

Woodland, J. M.; Barnett, C. J.; Dorman, D. E.;

Gruber, J. M.; Shih, C.; Spangle, L. A.; Wilson, T. M.; Ehlhardt, W. J.

CORPORATE SOURCE:.

Lilly Res. Laboratories, USA

SOURCE:

Drug Metabolism and Disposition (1997),

25(6), 693-700

CODEN: DMDSAI; ISSN: 0090-9556

PUBLISHER:

Williams & Wilkins

DOCUMENT TYPE:

LANGUAGE:

Journal English

The metabolism and disposition of LY231514 was studied in mice and dogs. LY231514 is a novel pyrrolopyrimidine-based multi-target antifolate (MTA) showing broad in vivo antitumor activity in mouse models and is currently in phase II human clin. trials. Doses (i.v.) of the compound showed high plasma levels, resulting in AUC values of 30-33 μg -hr/mL for mice and dogs after 20 and 7.5 mg/kg doses, resp. The compound was eliminated rapidly. Half-life values for mice and dogs were about 7 and 2 h, resp. In vitro plasma binding measured 56% in mice, 46% in dogs, and 81% in humans. Fecal elimination was the major excretion pathway in mice after single i.v. doses of [14C]LY231514. Urine constituted the major route of excretion in dogs. Parent LY231514 accounted for the majority of urinary radiocarbon in mice (90%) and dogs (68%). Minor metabolites were found in urine, but the amts. were too small to isolate or identify. Based on an earlier observation that LY231514 photodegraded to produce reaction products having similar retention times as these minor urinary isolates, a photo oxidation system was developed which in fact produced these

metabolites. Subsequently, these photolytically produced materials were used as stds. to identity two novel in vivo metabolites formed by oxidation of the pyrrolo-pyrimidine ring system of LY231514. The oxidative transformations are similar to those observed for tryptophan and other indoles in that the pyrrole ring is oxidized to give an amide; further oxidation cleaves this ring, one ring carbon is lost, and a ketone is formed. 193265-49-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(antifolate drug LY231514 metabolism and pharmacokinetics in mice and dogs) 193265-49-5 CAPLUS

RN 193265-49-5 CAPLUS CN Benzenebutanoic acid, α -(dicyanomethyl)-4-[(1,1-dimethylethoxy)carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ &$$

IT

L6 ANSWER 17 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1997:330734 CAPLUS

DOCUMENT NUMBER: 127:34293

TITLE: The reactions of Wittig-Horner reagents with

1,3-dioxo- Δ 2, α -indanmalononitrile

AUTHOR(S): Boulos, Leila Sadek; Yakout, El-Sayed M. A.

CORPORATE SOURCE: National Research Centre, Cairo, Egypt
SOURCE: Heteroatom Chemistry (1997), 8(3), 253-257

CODEN: HETCE8; ISSN: 1042-7163

PUBLISHER: Wiley
DOCUMENT TYPE: Journal
LANGUAGE: English

LANGUAGE: English

AB Wittig-Horner reagents react with 1,3-dioxo- $\Delta 2$, α indanmalononitrile to give phosphonate adducts. Structural reasoning for the new products was based on compatible anal. and spectral data (IR, 1H, 31P NMR, and $\dot{M}S$). The mechanism that accounts for the formation of the

new adducts is discussed. IT 190722-21-5P 190722-23-7P 190722-25-9P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 190722-21-5 CAPLUS

CN Butanedioic acid, 2-[dicyano(3-hydroxy-1-oxo-1H-inden-2-yl)methyl]-3-(diethoxyphosphinyl)-, dimethyl ester (9CI) (CA INDEX NAME)

RN 190722-23-7 CAPLUS

CN Butanedioic acid, 2-[dicyano(3-hydroxy-1-oxo-1H-inden-2-yl)methyl]-3-(diethoxyphosphinyl)-, diethyl ester (9CI) (CA INDEX NAME)

RN 190722-25-9 CAPLUS

REFERENCE COUNT:

THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 18 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

18

ACCESSION NUMBER:

1997:274825 CAPLUS

DOCUMENT NUMBER:

126:317775

TITLE:

Ring-opening polymerization of cyclobutane adduct of dimethyl 1,1-dicyanoethylene-2,2-dicarboxylate and

ethyl vinyl ether

AUTHOR(S):

Yokozawa, Tsutomu; Wakabayashi, Yuki; Kimura, Takamasa

CORPORATE SOURCE:

Dep. Applied Chem., Kanagawa Univ., Yokohama, 221,

Japan

SOURCE:

Journal of Polymer Science, Part A: Polymer Chemistry

(1997), 35(8), 1563-1570

CODEN: JPACEC; ISSN: 0887-624X

PUBLISHER:

Wiley Journal English

DOCUMENT TYPE: LANGUAGE:

AB For an extension of the work on the ring-opening polymns. of cyclobutane adducts of strong donor olefins and strong acceptor olefins yielding novel alternating copolymers of those olefins, the ring-opening polymerization of the cyclobutane adduct (I; di-Me 2,2-dicyano-3-ethoxy-1,1-cyclobutanedicarboxylate) of di-Me 1,1-dicyanoethylene-2,2-dicarboxylate (DDED) and Et vinyl ether (EVE) is investigated. I reacted with methanol and acetic acid at ambient temperature to yield the corresponding ring-opened adducts. Polymns. of I were carried out with anionic initiators, tertiary amines, ammonium halides, and Lewis acids, resp., according to the

polymerization

methods of the cyclobutane adduct of tetracyanoethylene and EVE. All these polymerization catalysts except for ammonium halides were effective for the

polymerization of I, yielding alternating copolymers of DDED and EVE. The chain

transfer reactions of the polymerization with anionic initiators are also discussed on the basis of a model reaction.

IT 189348-52-5

RL: RCT (Reactant); RACT (Reactant or reagent)

(model reaction for determination of mechanism; ring-opening polymerization of $\operatorname{di-Me}$

2,2-dicyano-3-ethoxy-1,1-cyclobutanedicarboxylate)

RN 189348-52-5 CAPLUS

CN Propanedioic acid, (dicyanomethyl)(2,2-diethoxyethyl)-, dimethyl ester (9CI) (CA INDEX NAME)

IT 189348-50-3P 189348-51-4P

RL: SPN (Synthetic preparation); PREP (Preparation)

(model reactions for polymerization; reactions of di-Me

2,2-dicyano-3-ethoxy-

1,1-cyclobutanedicarboxylate with acetic acid and methanol)

RN 189348-50-3 CAPLUS

CN Propanedioic acid, (dicyanomethyl)(2-ethoxy-2-methoxyethyl)-, dimethyl ester (9CI) (CA INDEX NAME)

RN- 189348-51-4 CAPLUS

CN Propanedioic acid, [2-(acetyloxy)-2-ethoxyethyl](dicyanomethyl)-, dimethyl ester (9CI) (CA INDEX NAME)

A STATE OF BUILDING

189348-53-6P IT

RL: SPN (Synthetic preparation); PREP (Preparation)

(ring-opening polymerization of di-Me 2,2-dicyano-3-ethoxy-1,1cyclobutanedicarboxylate)

RN 189348-53-6 CAPLUS

5-Octene-2,2,6-tricarboxylic acid, 1,1,5-tricyano-4,8,8-triethoxy-, CN trimethyl ester (9CI) (CA INDEX NAME)

27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 19 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1996:672685 CAPLUS

DOCUMENT NUMBER:

126:8777

TITLE:

Ring-Opening Polymerization of the Cyclobutane Adduct

of Methyl Tricyanoethylenecarboxylate and Ethyl Vinyl

AUTHOR(S):

Yokozawa, Tsutomu; Tsuruta, Ei-ichi

CORPORATE SOURCE:

Department of Applied Chemistry, Kanagawa University,

Yokohama, 221, Japan

SOURCE:

Macromolecules (1996), 29(25), 8053-8056 CODEN: MAMOBX; ISSN: 0024-9297

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE! English

The ring-opening polymns. of a cyclobutane adduct (I) of Me tricyanoethylenecarboxylate (MTCE) and Et vinyl ether (EVE) are investigated. The adduct I reacted with acetic acid and ethanol at ambient temperature to yield the ring-opened corresponding adducts in good yields. I was polymerized with Lewis acids, anionic initiators, tertiary amines, and ammonium halides. All the catalysts except for ammonium halides were effective for the alternating polymerization similar to the polymerization

of cyclobutane adduct of TCNE and EVE.

184092-92-0P 184092-93-1P IT

> RL: SPN (Synthetic preparation); PREP (Preparation) (ring-opening reactivity of cyclobutane adduct of Me

tricyanoethylenecarboxylate with ethanol or acetic acid and Et vinyl ether)

184092-92-0 CAPLUS RN

CN Butanoic acid, 4-(acetyloxy)-2-cyano-2-(dicyanomethyl)-4-ethoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 184092-93-1 CAPLUS

CN Butanoic acid, 2-cyano-2-(dicyanomethyl)-4,4-diethoxy-, methyl ester (9CI) (CA INDEX NAME)

9 REFERENCE COUNT: THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 20 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1996:664756 CAPLUS

DOCUMENT NUMBER:

125:329472

TITLE:

Preparation of ring-fused pyrimidine-containing amino

acid derivatives as antiprotozoan agents

INVENTOR(S):

Horii, Toshihiro; Aono, Tetsuya

PATENT ASSIGNEE(S):

Takeda Chemical Industries Ltd, Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 31 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	DATE			
JP 08225574	A ·	19960903	JP 1995-330939	19951220 <		
PRIORITY APPLN. INFO.:			JP 1995-330939 A	19951220		
			JP 1994-317938	19941221		

OTHER SOURCE(S): MARPAT 125:329472

For diagram(s), see printed CA Issue.

The title compds. [I; ring A = (un)substituted five membered ring; Z = AB (un) substituted bivalent aliphatic chain consisting of a series of <5 atoms and optionally interrupted by one hetero atom in the chain; B = (un) substituted 5- or 6-membered heterocyclyl or carbocyclyl; the substituent of B is preferably CONHCH(CO2R3)(CH2)pWR4; wherein p = 1-4; W = bond, O, NHCONH, NR, NRCO, CONR, NHSO2; wherein R = H, C1-4 hydrocarbyl; CO2R3 = optionally esterified CO2H; R4 = (un)substituted chain or cyclic group; or Z = (CR1R2)n-Z1; wherein R1, R2 = H,lower alkyl; Z1 = bond, O,

NH; n = 1-5] or salts thereof, which are useful for treating infections of protozoa, particularly coccidium and drug-resistant malaria, are prepared Thus, 4-[2-(2,4-diamino-7H-pyrrolo[2,3-d]pyrimidin-5-yl)ethyl]benzoic acid ditrifluoroacetate was condensed with Me O-(4-methoxycarbonylbenzyl)-L-serinate hydrochloride using di-Et cyanophosphate and Et3N in DMF at room temperature for 1 h to give 66% Me N-[4-[2-(2,4-diamino-7H-pyrrolo[2,3-d]pyrimidin-5-yl)ethyl]benzoyl]-O-O-(4-methoxycarbonylbenzyl)-L-serinate, which was saponified with a mixture of 1 h aqueous NaOH and MeOH at room temperature for

5 h and neutralized with dilute HCl to give 84% N-[4-[2-(2,4-diamino-7H-pyrrolo[2,3-d]pyrimidin-5-yl)ethyl]benzoyl]-0-0-(4-methoxycarbonylbenzyl)-L-serine. These compds. I inhibited dihydrofolic acid reductase of malaria protozoa Plasmodium falciparum with IC50 of 0.8-62 nM. The title compound (II; R5 = Q) showed ED50 of 0.17 and 0.11 nM for inhibiting the proliferation of wild type-malaria protozoa P. falciparum 3D7 and cycloguanyl-resistant P. falciparum FCR3, resp. Capsule, tablet, and vial formulations containing II (R5 = Q1) were prepared 182961-44-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of ring-fused pyrimidine-containing amino acid derivs. as antiprotozoan agents)

RN 182961-44-0 CAPLUS

IT

CN Benzenepentanoic acid, α -(dicyanomethyl)-3,4,5-trimethoxy-, ethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 21 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1996:365832 CAPLUS

DOCUMENT NUMBER: 125:86845

TITLE: Cyclopropenation and Related Reactions of Ruthenium

Vinylidene Complexes

AUTHOR(S): Ting, Pei-Chen; Lin, Ying-Chih; Lee, Gene-Hsiang;

Cheng, Ming-Chu; Wang, Yu

CORPORATE SOURCE: Department of Chemistry, National Taiwan University,

Taipei, 106, Taiwan

SOURCE: Journal of the American Chemical Society (1996

), 118(27), 6433-6444

CODEN: JACSAT; ISSN: 0002-7863

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

AB Facile deprotonation of a number of cationic ruthenium vinylidene complexes, followed by cyclopropenation, is accomplished in acetone. The deprotonation of [Ru]:C:(Ph)CH2R+, ([Ru] = (η5-C5H5)(PPh3)2Ru through out this abstract) by n-Bu4NOH induces a novel cyclization reaction and yields the neutral cyclopropenyl complexes [cyclic] [Ru]-C:C(Ph)CHR (3b, R = CN; 3c, R = Ph; 3d, R = CH:CH2; 3e, R = CH:CMe2). Cyclic complex [Ru]-C:C(C6H9)CHCN+ is similarly prepared Protonation of 3b-3e regenerates

the corresponding vinylidene complexes. Deprotonation of [Ru]:C:C(Ph)CH2COOMe+ by n-Bu4NOH induces a different type of cyclization and yields the neutral furan complex [cyclic] [Ru]-C:C(Ph)CH:C(O)OMe (4h). The cyclopropenyl complex containing a methoxy substituent cannot be prepared from [Ru]:C:C(Ph)CH2OCH3+ (2i), but F- of n-Bu4NF attacks the Ca of 2i to produce the unstable vinyl complex [Ru]C(F):C(Ph)CH2OCH3. Cyclic complex [Ru]-C:C(Ph)C(CN)OCH3 (9b) was indirectly prepared from the addition of TCNQ to 3b, giving [cyclic] [Ru]:C:C(Ph)CH(CN)TCNQ (6b) followed by methanolysis. Unlike 3, complex 9b is not converted to vinylidene complex, instead, removal of the methoxy substituent by acid gives the cationic cyclopropenylium complex [Ru]-C:C(Ph)C(CN)+. Cyclic complex [Ru]-C:C(Ph)C(COOMe)+ is similarly prepared from 4h via a TCNQ complex followed by a methoxy-substituted complex. In the presence of allyl iodide, opening of the three-membered ring of 3b, followed by a subsequent oxidative coupling reaction, gives a dimeric dicationic product {[Ru]:C:C(Ph)-CHCN}22+ (11). Proton abstraction of 11 by n-Bu4NOH gives the biscyclopropenyl complex {[Ru]-C:C(Ph)CCN}2. Mol. structures of complexes 3b, 4h, 6b, 9b, 11, and [cyclic] [Ru]-C:C(Ph)C(CPh3)CN have been confirmed by x-ray diffraction anal.

IT 178687-62-2P

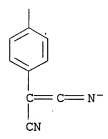
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(cyclopropenation and related reactions of ruthenium vinylidene complexes)

RN 178687-62-2 CAPLUS

CN Ruthenium, (n5-2,4-cyclopentadien-1-yl)[4,4-dicyano-4-[4-(cyanoiminatoethenyl)phenyl]-3-(methoxycarbonyl)-2-phenyl-1-butenylidene]bis(triphenylphosphine)- (9CI) (CA INDEX NAME)

PAGE 1-A



ANSWER 22 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1996:6671 CAPLUS

DOCUMENT NUMBER:

124:177091

TITLE:

Novel gem-dinitrile functionalized polyesters and

polyamides from malononitrile; potential piezoelectric

materials

AUTHOR(S):

CÓRPORATE SOURCE:

Steadman, Scott; Parrish, Dennis A.; Mathias, Lon J. Department of Polymer Science, University of Southern

Mississippi, Hattiesburg, MS, 39406-0076, USA

SOURCE:

Polymer Preprints (American Chemical Society, Division

of Polymer Chemistry) (1995), 36(2), 320-1

CODEN: ACPPAY; ISSN: 0032-3934

PUBLISHER:

American Chemical Society, Division of Polymer

Chemistry

DOCUMENT TYPE:

Journal

LANGUAGE: English

Novel polyamides and polyesters in which the dinitrile group can potentially align in the same direction as the dipole of the carbonyl groups were synthesized via step growth dialkylation of malononitrile. Diamide and diester monomers facilitated polymerization by the attachment of chlorine to an activated position (α to carbonyl). The polymers, having mol. weight 5000-8000, were characterized by NMR, viscosity and thermal anal.

IT 169893-85-0P 174297-80-4P 174297-82-6P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation of gem-dinitrile functionalized polyesters and polyamides from malononitrile as potential piezoelec. materials)

169893-85-0 CAPLUS RN

CN Poly[oxy-1,3-propanediyloxy(3,3-dicyano-1,5-dioxo-1,5-pentanediyl)] (9CI) (CA INDEX NAME)

RN 174297-80-4 CAPLUS

CN Poly[oxy-1,4-butanediyloxy(3,3-dicyano-1,5-dioxo-1,5-pentanediyl)] (9CI) (CA INDEX NAME)

RN 174297-82-6 CAPLUS
CN Poly[oxy-1,4-cyclohexanediyloxy(3,3-dicyano-1,5-dioxo-1,5-pentanediyl)]
(9CI) (CA INDEX NAME)

L6 ANSWER 23 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1995:900662 CAPLUS

DOCUMENT NUMBER: 124:116317

TITLE: Lanthanum isopropoxide catalyzed addition of activated

nucleophiles to imines

AUTHOR(S): Yamamoto, Yoshinori; Fukui, Hiroyuki; Honda, Yoshihiro

CORPORATE SOURCE: Dept. Chem., Tohoku Univ., Sendai, 980-77, Japan

SOURCE: Applied Organometallic Chemistry (1995), 9(5

& 6), 467-71

CODEN: AOCHEX; ISSN: 0268-2605

PUBLISHER: Wiley
DOCUMENT TYPE: Journal
LANGUAGE: English

OTHER SOURCE(S): CASREACT 124:116317

AB The addition of certain activated nucleophiles to activated imines is catalyzed by lanthanum isopropoxide. As activated nucleophiles, methylmaloninitrile and Me 2-cyanopropanoate can be utilized. Imines having an electron-withdrawing group either at the carbon or at the nitrogen atom of the C:N double bond can be used: for example N-toluenesulfonylimines, N-(4-methoxycarbonylphenyl)imines and α -imino esters.

IT 155751-02-3P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (lanthanum isopropoxide catalyzed addition of activated nucleophiles to imines)

RN 155751-02-3 CAPLUS

CN Butanoic acid, 3,3-dicyano-2-[(1-phenylethyl)amino]-, 5-methyl-2-(1-methyl-1-phenylethyl)cyclohexyl ester, $[1R-[1\alpha[S*(S*)],2\beta,5\alpha]]-(9CI)$ (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

(9CI) (CA INDEX NAME)

RN 155696-72-3 CAPLUS CN Butanoic acid, 3,3-dicyano-2-[(1-phenylethyl)amino]-, 5-methyl-2-(1-methyl-1-phenylethyl)cyclohexyl ester, [1R-[1 α [S*(R*)],2 β ,5 α]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 172880-55-6 CAPLUS

CN Butanoic acid, 3,3-dicyano-2-[(1-phenylethyl)amino]-, butyl ester, (R*,R*)- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 172880-56-7 CAPLUS

CN Butanoic acid, 3,3-dicyano-2-[(1-phenylethyl)amino]-, butyl ester, (R*,S*)- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 173006-24-1 CAPLUS

CN Butanoic acid, 3,3-dicyano-2-[(1-methylethyl)amino]-, 5-methyl-2-(1-methyl-1-phenylethyl)cyclohexyl ester, [1R-[1 α (R*),2 β ,5 α]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 173006-25-2 CAPLUS

CN Butanoic acid, 3,3-dicyano-2-[(1-phenylethyl)amino]-, 5-methyl-2-(1-methyl-1-phenylethyl)cyclohexyl ester, $[1R-[1\alpha[R^*(R^*)],2\beta,5\alpha]]-(9CI)$ (CA INDEX NAME)

Absolute stereochemistry.

RN 173006-26-3 CAPLUS

CN Butanoic acid, 3,3-dicyano-2-[(1-phenylethyl)amino]-, 5-methyl-2-(1-methyl-1-phenylethyl)cyclohexyl ester, $[1R-[1\alpha[R^*(S^*)],2\beta,5\alpha]]-(9CI)$ (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 24 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1995:845626 CAPLUS

DOCUMENT NUMBER: 124:86769

TITLE: Novel three-component reaction of 1,1-dicyano-2-

(trifluoromethyl)ethylenes with primary arylamines and

ketones

AUTHOR(S): Tyutin, V. Yu.; Chkanikov, N. D.; Nesterov, V. N.;

Antipin, M. Yu.; Struchkov, Yu. T.; Kolomiets, A. F.;

Fokin, A. V.

CORPORATE SOURCE: A. N. Nesmeyanov Inst. Organoelem. Compd., Russ. Acad.

Sci., Moscow, 117813, Russia

SOURCE:

Izvestiya Akademii Nauk, Seriya Khimicheskaya (

1993), (3), 552-9 CODEN: IASKEA

PUBLISHER:

Nauka Journal

DOCUMENT TYPE:

Russian

LANGUAGE:

CASREACT 124:86769

OTHER SOURCE(S):

1,1-Dicyano-2,2-bis(trifluoromethyl)ethylene and 3,3-dicyano-2-(trifluoromethyl)acrylates react with primary arylamines in the presence of ketones to form 1,1-aryl-1,4-dihydropyridine derivs. under mild conditions. In this three-component reaction Schiff bases are formed as intermediates. 1,4-Dihydropyridines derivs., which are the products of

intermediates. 1,4-Dihydropyridines derivs., which are the products of three-component heterocyclization, were also obtained by interaction of the corresponding Schiff bases with 1,1-dicyano-2-

(trifluoromethyl) ethylenes.

IT 134641-39-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 134641-39-7 CAPLUS

CN Benzeneacetic acid, 4-amino- α -(dicyanomethyl)-3,5-dimethyl- α -

(trifluoromethyl) -, ethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 25 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1995:695727 CAPLUS

DOCUMENT NUMBER:

123:286966

TITLE:

Novel gem-dinitrile functionalized polyesters and

polyamides from malononitrile; potential piezoelectric

materials

AUTHOR(S):

Mathias, Lon J.; Parrish, Dennis A.; Steadman, Scott

CORPORATE SOURCE:

Department Polymer Science, University Southern

Mississippi, Hattiesburg, MS, 39406-0076, USA

SOURCE:

Polymer Preprints (American Chemical Society, Division

of Polymer Chemistry) (1994), 35(2), 659-60

CODEN: ACPPAY; ISSN: 0032-3934

PUBLISHER:

American Chemical Society, Division of Polymer

Chemistry

DOCUMENT TYPE:

LANGUAGE:

Journal English

AB The initial success is described in obtaining a polyester and polyamide in which the dinitrile group net dipole can potentially align in the same direction as the carbonyl groups.

IT 169893-85-0P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of gem-dinitrile functionalized polyesters and polyamides from malononitrile as potential piezoelec. materials)

RN 169893-85-0 CAPLUS

CN Poly[oxy-1,3-propanediyloxy(3,3-dicyano-1,5-dioxo-1,5-pentanediyl)] (9CI) (CA INDEX NAME)

L6 ANSWER 26 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1995:440268 CAPLUS

DOCUMENT NUMBER:

123:112653

TITLE:

Synthesis and antitumor activity of

pyrrolo[2,3-d]pyrimidine antifolates with a bridge

chain containing a nitrogen atom

AUTHOR(S):

Aso, Kazuyoshi; Hitaka, Takenori; Yukishige, Koichi;

Ootsu, Koichiro; Akimoto, Hiroshi

CORPORATE SOURCE:

Pharmaceutical Res. Div., Takeda Chem. Industries,

Ltd., Osaka, 532, Japan

SOURCE:

Chemical & Pharmaceutical Bulletin (1995),

43(2), 256-61

CODEN: CPBTAL; ISSN: 0009-2363 Pharmaceutical Society of Japan

PUBLISHER: DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 123:112653

GI

AB Novel pyrrolo[2,3-d]pyrimidine antifolates I (R = H, Me) with a nitrogen atom in the bridge chain between the 2,4-diaminopyrrolo[2,3-d]pyrimidine and phenylene rings were designed and efficiently synthesized. I exhibited more potent inhibitory activities than methotrexate (MTX) against the proliferation of human epidermoid carcinoma KB cells and human non-small cell lung carcinoma A549 cells despite their modest dihydrofolate reductase (DHFR)-inhibitory potency.

IT 133719-38-7P 133719-41-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and antitumor activity of pyrrolo[2,3-d]pyrimidine antifolates with nitrogen-containing bridge chains)

RN 133719-38-7 CAPLUS

CN Benzoic acid, 4-[[4,4-dicyano-3-(methoxycarbonyl)butyl]methylamino]-, ethyl ester (9CI) (CA INDEX NAME)

RN 133719-41-2 CAPLUS

CN Benzoic acid, 4-[[4,4-dicyano-3-(ethoxycarbonyl)butyl][(1,1dimethylethoxy)carbonyl]amino]-, ethyl ester (9CI) (CA INDEX NAME)

ANSWER 27 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1994:434408 CAPLUS

DOCUMENT NUMBER:

121:34408

TITLE:

Transition metal catalyzed addition of certain

nucleophiles to imines

AUTHOR(S):

Yamamoto, Yoshinori; Kubota, Yasufumi; Honda,

Yoshihiro; Fukui, Hiroyuki; Asao, Naoki; Nemoto, Hisao

CORPORATE SOURCE:

Faculty of Science, Tohoku University, Sendai, 980,

Japan

SOURCE:

Journal of the American Chemical Society (1994

), 116(7), 3161-2

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE:

LANGUAGE:

Journal English

OTHER SOURCE(S):

CASREACT 121:34408

AB Imines react with certain nucleophiles in the presence of catalytic amts. of transition metal complexes to give alkylation products in good yield. Thus, imine (I) was treated with CH(CN)2(CO2Et) in the presence of RhHCO(PPh3)3 in THF solvent to give alkylation product II in 75% yield. A significantly high diastereomeric excess was accomplished by using III [R1 = (-)-8-phenylmenthyl] in which a chiral auxiliary exists at the ester unit. The Ls(O-iso-Pr)3 catalyzed reaction of III with CH(CN)2Me in THF at room temperature gave IV (R1 as above) as the predominant diastereoisomer in a 90:10 ratio; x-ray anal. indicate that the α -carbon to the amino group possesses the S configuration.

IT 155751-02-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, absolute configuration of)

RN 155751-02-3 CAPLUS

CN Butanoic acid, 3,3-dicyano-2-[(1-phenylethyl)amino]-, 5-methyl-2-(1-methyl-1-phenylethyl)cyclohexyl ester, $[1R-[1\alpha[S^*(S^*)],2\beta,5\alpha]]-(9CI)$ (CA INDEX NAME)

Absolute stereochemistry.

IT 155696-71-2P 155696-72-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, stereoselective)

RN 155696-71-2 CAPLUS

CN Butanoic acid, 3,3-dicyano-2-[(1-methylethyl)amino]-, 5-methyl-2-(1-methyl-

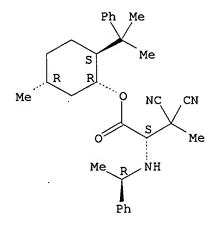
1-phenylethyl)cyclohexyl ester, [1R-[1 α (S*),2 β ,5 α]]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 155696-72-3 CAPLUS

CN Butanoic acid, 3,3-dicyano-2-[(1-phenylethyl)amino]-, 5-methyl-2-(1-methyl-1-phenylethyl)cyclohexyl ester, [1R-[1 α [S*(R*)],2 β ,5 α]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



L6 ANSWER 28 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1994:269808 CAPLUS

DOCUMENT NUMBER: 120:269808

TITLE: Wittig reactions of a fluoren-9-ylidene and an

anthrone-10-arylidene

AUTHOR(S): Ganoub, Neven A. F.

CORPORATE SOURCE: Dep. Pesticide Chem., Natl. Res. Cent., Cairo, Egypt

SOURCE: Phosphorus, Sulfur and Silicon and the Related

Elements (1993), 81(1-4), 125-31

CODEN: PSSLEC; ISSN: 1042-6507

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 120:269808

GΙ

AB The Wittig reactions of fluoren-9-ylidenemalonitrile (I) and 10-benzylideneanthrone (II) with phosphonium ylides Ph3P+CH-CO2R (R = Me, Et) have been investigated. In both cases, unusual reaction products, e.g., bis(9-fluorenyl)cyclopentane III (from I), were isolated and identified on the basis of elemental analyses and spectral studies.

IT 154496-99-8P 154497-00-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and thermal decomposition of)

RN 154496-99-8 CAPLUS

CN 9H-Fluorene-9-propanoic acid, β , β -dicyano- α - (triphenylphosphoranylidene)-, methyl ester (9CI) (CA INDEX NAME)

RN 154497-00-4 CAPLUS
CN 9H-Fluorene-9-propanoic ac

9H-Fluorene-9-propanoic acid, β , β -dicyano- α - (triphenylphosphoranylidene)-, ethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 29 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1994:244629 CAPLUS

DOCUMENT NUMBER: 120:244629

TITLE: Synthesis of spiro indolin-2-one derivatives

AUTHOR(S):

El-Ahl, Abdel Aziz S.; Afeefy, Hussein; Metwally,

Mohamed Abbas

CORPORATE SOURCE:

SOURCE:

Fac. Sci., Mansoura Univ., Mansoura, Egypt Journal of Chemical Research, Synopses (1994

), (1), 14-15

CODEN: JRPSDC; ISSN: 0308-2342

DOCUMENT TYPE:

LANGUAGE:

Journal -English

OTHER SOURCE(S):

CASREACT 120:244629

GI

AΒ Title compds. I were prepared by heating dicyanomethyleneindolinones II (R = H, R1 = H, Me; R = Me, Ac, R1 = H) with active methylene compds., XCH2COZ (X = Ac, Z = OEt, Me; X = cyano, Z = Ph).

154379-70-1P 154379-71-2P IT

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

154379-70-1 CAPLUS RN

Benzeneacetic acid, 2-(acetylamino)- α -(dicyanomethyl)-, methyl ester CN (CA INDEX NAME)

RN 154379-71-2 CAPLUS

Benzeneacetic acid, 2-(acetylamino)- α -(dicyanomethyl)-, ethyl ester CN (9CI) (CA INDEX NAME)

L6 ANSWER 30 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1993:213100 CAPLUS

DOCUMENT NUMBER: 118:213100

TITLE: Preparation of tricyclic fused pyrimidine compounds

INVENTOR(S): Akimoto, Hiroshi; Otsu, Koichiro; Miwa, Tetsuo

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 25 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE			
JP 04211063	Α	19920803	JP 1991-65613	19910305 <			
PRIORITY APPLN. INFO.:			JP 1990-54620 A1	19900305			
OTHER SOURCE(S):	MARPAT	118:213100		•			

OTHER SOURCE(S): MARPAT 118:213100
GI For diagram(s), see printed CA Issue.

The title compds. [I; Q1 = H, halo, radical linked through C, N, O, or S; one of Q2 and Q3 = N, the other = N, CH; Y = N, CR1 (wherein R1 = H, hydrocarbyl), methylidyne; Z = C2-5 bivalent radical containing optional substituents; ring A1, A2 = (substituted) 5-7-membered ring; B = (substituted) cyclic radical, etc.], useful as antitumor agents with high selectivity, are prepared Cyclocondensation of 1.181 g ester II (preparation given) with 314 mg guanidine HCl and Me3COK in Me3COH gave 1.02 g pyrrolopyrimidine III, which (1.010 g) was treated with borane-THF complex in THF at 0° and then at 50°, the solution cooled and stirred with HOAC-MeOH at room temperature to give 542 mg IV. The preferred deads of I

with HOAc-MeOH at room temperature to give 542 mg IV. The preferred doses of I are 2.0-500 mg/kg-day orally and 1.0-200 mg/kg injection.

IT 147239-87-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, in preparation of antitumor agent)

RN 147239-87-0 CAPLUS

CN

Benzenepentanoic acid, α -(dicyanomethyl)- β -(dimethoxymethyl)-4-[(1,1-dimethylethoxy)carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 31 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1993:124655 CAPLUS

DOCUMENT NUMBER: 118:124655

TITLE: Wittig reaction of 1-(dicyanomethylene)acenaphthen-2-

one

AUTHOR(S): Abdou, Wafaa M.; Ganoub, Neven A. F.

CORPORATE SOURCE: Natl. Res. Cent., Cairo, Egypt

SOURCE: Heteroatom Chemistry (1992), 3(2), 133-7

CODEN: HETCE8; ISSN: 1042-7163

DOCUMENT TYPE:

LANGUAGE:
OTHER SOURCE(S):

Journal English

CASREACT 118:124655

GΙ

AB The Wittig reactions of title compound I with alkoxycarbonylmethylenetriphen ylphosphoranes Ph3P+C-HCOR (R = OMe, OEt) were investigated and the reaction products zwitterion II and heterocycles III and IV were isolated. Reaction of I with benzoylmethylenetriphenylphosphorane Ph3P+C-HCOPh proceeded only at high temperature, yielding V and III (R = Ph). Mechanisms accounting for the formation of the adducts are discussed. Wittig olefination of several products was studied.

IT 145882-80-0P 145882-83-3P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 145882-80-0 CAPLUS

CN 1-Acenaphthylenepropanoic acid, β , β -dicyano-1, 2-dihydro-2-oxo- α -(triphenylphosphoranylidene)-, methyl ester (9CI) (CA INDEX NAME)

RN 145882-83-3 CAPLUS

CN 1-Acenaphthylenepropanoic acid, β , β -dicyano-1, 2-dihydro-2-oxo- α -(phenylmethylene) - (9CI) (CA INDEX NAME)

IT 145882-82-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation, Wittig olefination, and thermal intramol. cyclocondensation of)

RN 145882-82-2 CAPLUS

CN Phosphonium, [2,2-dicyano-2-(2-hydroxy-1-acenaphthylenyl)-1-(methoxycarbonyl)ethyl]triphenyl-, inner salt (9CI) (CA INDEX NAME)

L6 ANSWER 32 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1993:29863 CAPLUS

DOCUMENT NUMBER:

118:29863

TITLE:

Silver halide photographic material containing a

compound which releases photographically useful

species upon development

INVENTOR(S):

Asatake, Atsushi

PATENT ASSIGNEE(S): SOURCE:

Konica Co., Japan Jpn. Kokai Tokkyo Koho, 19 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	ŧ
JP 04177243	Α	19920624	JP 1990-305540	19901110 <	
PRIORITY APPLN. INFO.:			JP 1990-305540	19901110	
AB The photog. materi	ial conta	ains a compo	ound CRR1R2C(R3R4)mZ(Z1)	nPUG (R =	
leaving group rele	eased by	nucleophili	ic substitution; R1,R2,F	R3,R4 = H,	

leaving group released by nucleophilic substitution; R1,R2,R3,R4 = H, aliphatic, aromatic, heterocyclic or electron-attracting group; Z = electron-attracting group; Z1 = timing group to be subjected to break and release PUG: PUG = photog. useful group; m,n = 0, 1). The photog. material has good storage stability, while upon development, it releases the PUGs at a proper reaction rate even in developer solution of relatively low pH.

IT 144896-71-9 145059-42-3

RL: USES (Uses)

(photog. useful group-releasing, in processing)

RN 144896-71-9 CAPLUS

CN Butanoic acid, 4-[[4-(3-chloro-3,3-dicyano-1-oxopropoxy)-3-[[(5-methyl-1,3,4-oxadiazol-2-yl)thio]methyl]phenyl]amino]-4-oxo- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & & & \\ & & & & \\ NH-C-CH_2-CH_2-CO_2H \\ \hline & & & \\ NH-C-CH_2-CH_2-CO_2H \\ \hline & & & \\ CN & & & \\ CN & & & \\ Me & & & \\ Me & & & \\ \end{array}$$

RN 145059-42-3 CAPLUS

Cl

CN Dodecanoic acid, 2-(chlorodicyanomethyl)-, 4-[[2-[[4-[[1,5-dihydro-3-methyl-5-oxo-1-(4-sulfophenyl)-4H-pyrazol-4-ylidene]methyl]-3-methylphenyl]methylamino]ethoxy]methyl]-3-methyl-1-(4-nitrophenyl)-1H-pyrazol-5-yl ester, monopotassium salt (9CI) (CA INDEX NAME)

PAGE 1-A

$$NC-C-CN$$
 $Me-(CH_2) g-CH-C-O$
 $N-CH_2-O-CH_2-CH_2-N$
 Me
 Me

L6 ANSWER 33 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1992:174018 CAPLUS

DOCUMENT NUMBER:

116:174018

TITLE:

Synthesis and structure-antimicrobial activity

relationships of quaternary ammonium derivatives of

perhydropyrrolo-[3,4-c]pyridine

AUTHOR(S):

Altomare, C.; Carotti, A.; Casini, G.; Cellamare, S.;

Ferappi, M.; Vitali, C.

CORPORATE SOURCE:

Dip. Farm. Chim., Univ. Bari, Bari, I-70125, Italy

SOURCE:

Arzneimittel-Forschung (1992), 42(2), 152-5

CODEN: ARZNAD; ISSN: 0004-4172

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 116:174018

GI

AB A homologous series of perhydropyrrolo[3,4-c]pyridine quaternary ammonium derivs. I (n = 5, 7, 9-13,15) was synthesized from EtO2CCH2CH(CO2Et)CH(CN)2 and tested for in vitro antibacterial activity against different gram-pos. and gram-neg. bacteria. All I were more potent than the reference compound, benzalkonium chloride. Antibacterial activity, expressed as log 1/MIC, was linearly related to lipophilicity up to C13-C14 homologs, where a break in the linear relationship was observed

IT 82584-86-9

RL: RCT (Reactant); RACT (Reactant or reagent)
 (hydrogenation-intramol. cyclocondensation of)

RN 82584-86-9 CAPLUS

CN Butanedioic acid, (dicyanomethyl)-, diethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 34 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1992:83465 CAPLUS

DOCUMENT NUMBER:

116:83465

TITLE:

The regioselectivity of the ring opening of 1-activated or nonactivated 2-alkoxycarbonyl or 2-cyanoaziridines by carbanions of the dicarbonyl

compounds

AUTHOR(S):

Bouayad, Zoheir; Chanet-Ray, Josette; Ducher, S.;

Vessiere, Roger

CORPORATE SOURCE:

Ec. Natl. Super. Chim. Clermont-Ferrand, Univ. Blaise

Pascal, Aubiere, 63177, Fr.

SOURCE:

Journal of Heterocyclic Chemistry (1991),

28(7), 1757-67

CODEN: JHTCAD; ISSN: 0022-152X

DOCUMENT TYPE:

Journal

LANGUAGE:

English

CO2CHMe2

AB Aziridines, e.g. I, reacted with carbanions of dicarbonyl compds., e.g. RO2CCH2CO2R (R = Me, Et, CHMe2), to give ring opened products and/or ring enlarged products, e.g. (RO2C)2CHCH2CH(NHBz)CO2CHMe2, (RO2C)2CHCH(CO2CHMe2)CH2NHBz, and pyrrole II. The regioselectivity depends on several factors. The Ph group on C-3 favors C-3-N bond cleavage, whereas C-2-N bond cleavage is predominant with C-3 substituted or C-2-H aziridines. Cyanoaziridines are predominantly cleaved at C-3-N.

II

IT 138478-35-0P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

N 138478-35-0 CAPLUS

CN Propanoic acid, 2-[(benzoylamino)methyl]-3,3-dicyano-, 1-methylethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 35 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1991:680104 CAPLUS

DOCUMENT NUMBER:

115:280104

TITLE:

E/Z isomerization, solvolysis, addition, and

cycloaddition reactions of (E)-tert-butylketene methyl

tert-butyldimethylsilyl acetal

AUTHOR(S):

Adam, Waldemar; Wang, Xiaoheng

CORPORATE SOURCE:

Inst. Org. Chem., Univ. Wuerzburg, Wuerzburg, D-8700,

Germany

SOURCE:

Journal of Organic Chemistry (1991), 56(26),

7244-50

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 115:280104

AB In the presence of catalytic amts. of CF3COMe or CF3COCF3, the silvl ketene acetal Me3CCH:C(OMe)OSiMe2CMe3 (E-I) was isomerized into its Z isomer (Z/E ratio 90:10). For this novel E/Z isomerization a mechanism is proposed in which addition and reelimination of the fluoro ketone through a 1,4-dipolar intermediate operates. With the protic nucleophiles MeOH, CF3CH2OH, or PhOH, the ketene acetal E-I afforded the ortho esters Me3CCH2C(OMe)(OR)OSiMe2CMe3 (R = Me, CF3CH2, Ph) as addition products, while AcOH, CF3CO2H, or H2O led to Me pivalate as the solvolysis product. This chemical is readily explained through protonation of the ketene acetal E-I to generate the corresponding carbenium ion. At low temperature the reaction with TCNE gave the silylketene imine as labile cycloadduct, which underwent desilylation on workup to give the TCNE-incorporated ester (NC)2CHC(CN)2CH(CMe2)C(0)OMe; the latter eliminated hydrogen cyanide at room temperature to give the ene ester. With MTAD the labile silyl ene product was obtained initially, which underwent silyl migration to give N-silylated urazole; final desilylation led to the stable urazole II. Also, for the ene reactions of TCNE and MTAD with the silyl ketene acetal E-I, intervention of a 1,4-dipolar intermediate is proposed.

IT 136911-64-3P

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and hydrogen cyanide elimination of)

RN136911-64-3 CAPLUS

CN Butanoic acid, 3,3,4,4-tetracyano-2-(1,1-dimethylethyl)-, methyl ester (CA INDEX NAME)

IT 136911-63-2P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 136911-63-2 CAPLUS

CN 4-Pentenoic acid, 3,3,4-tricyano-2-(1,1-dimethylethyl)-5-[[(1,1-

dimethylethyl)dimethylsilyl]imino]-, methyl ester (9CI) (CA INDEX NAME)

ANSWER 36 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1991:608100 CAPLUS

DOCUMENT NUMBER:

115:208100

TITLE:

Chemistry of phosphorus ylides. 10. Reaction with

phosphacumulenes. IV. Synthesis of pyran, phosphoranylidene, oxaphosphorin and oxazaphosphorin

from the reaction of 1,3-dioxo- Δ 2, α indanmalononitrile with phosphoranes and

iminophosphoranes

AUTHOR(S):

Soliman, Fouad M.; Said, Medhat M.

CORPORATE SOURCE:

Natl. Res. Cent., Cairo, Egypt

SOURCE:

Phosphorus, Sulfur and Silicon and the Related

Elements (1991), 61(3-4), 335-40 CODEN: PSSLEC; ISSN: 1042-6507

DOCUMENT TYPE:

LANGUAGE:

Journal English

OTHER SOURCE(S):

CASREACT 115:208100

GI

AB 1,3-Dioxo- Δ 2, α -indanmalononitrile (I) reacts with the active ketenylidene- and thioketenylidenetriphenylphosphoranes Ph3P:C:C:X (X = 0, S, resp.) to give the corresponding pyrans II (X = 0, S). The reaction of II with 4-O2NC6H4CHO proceeds according to the Wittig reaction to give the resp. methylidene derivs. On the other hand, phosphoranylidenes III (R = acyl, alkoxycarbonyl) were isolated from the reaction of stable phosphoranes Ph3P:CHR with I. Moreover, an oxaphosphorin and oxazaphosphorin were prepared from the reaction of I with the phosphorane Ph3P:CPh2 and the iminophosphorane Ph3P:NCO2Et, resp.

IT 136829-50-0P 136848-91-4P

RN 136829-50-0 CAPLUS

CN 1H-Indene-2-propanoic acid, β , β -dicyano-3-hydroxy-1-oxo- α - (triphenylphosphoranylidene)-, methyl ester (9CI) (CA INDEX NAME)

RN 136848-91-4 CAPLUS

CN 1H-Indene-2-propanoic acid, β , β -dicyano-3-hydroxy-1-oxo- α - (triphenylphosphoranylidene)-, ethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 37 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1991:492003 CAPLUS 115:92003

DOCUMENT NUMBER: TITLE:

C-Alkylation of indoles with 1,1-bis(trifluoromethyl)-

2,2-dicyanoethylene and 2-trifluoromethyl-3,3-

dicyanoacrylic acid esters

AUTHOR(S):

Chkanikov, N. D.; Komarov, K. V.; Tyutin, V. Yu.;

Kolomiets, A. F.; Fokin, A. V.

CORPORATE SOURCE:

Inst. Elementoorg. Soedin. im. Nesmeyanova, Moscow,

USSR

SOURCE:

Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (

1991), (5), 1193-5

CODEN: IASKA6; ISSN: 0002-3353

DOCUMENT TYPE:

Journal

LANGUAGE:

Russian

OTHER SOURCE(S):

CASREACT 115:92003

GΙ

AB The indole derivs. I (R = H, Me; R1 = H, Me, Ph) were alkylated with (NC)2C:CR2CF3 (R2 = CF3, CO2Me, CO2Et) to give the corresponding dicyanoethyl derivs. II.

II

IT 135578-14-2P 135578-15-3P 135578-17-5P 135578-18-6P 135578-19-7P 135578-20-0P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 135578-14-2 CAPLUS

CN lH-Indole-3-acetic acid, α -(dicyanomethyl)- α -(trifluoromethyl)-, methyl ester (9CI) (CA INDEX NAME)

RN 135578-15-3 CAPLUS

CN 1H-Indole-3-acetic acid, α -(dicyanomethyl)- α -(trifluoromethyl)-, ethyl ester (9CI) (CA INDEX NAME)

RN 135578-17-5 CAPLUS

CN 1H-Indole-3-acetic acid, α -(dicyanomethyl)-2-methyl- α - (trifluoromethyl)-, methyl ester (9CI) (CA INDEX NAME)

RN 135578-18-6 CAPLUS

CN 1H-Indole-3-acetic acid, α -(dicyanomethyl)-2-methyl- α -

$$\begin{array}{c|c} & & \text{Me} \\ \hline & & \text{Me} \\ \hline & & \text{CF3} \\ \hline & & \text{CH-CN} \\ \hline & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ \end{array}$$

RN 135578-19-7 CAPLUS

CN lH-Indole-3-acetic acid, α -(dicyanomethyl)-1-methyl-2-phenyl- α -(trifluoromethyl)-, methyl ester (9CI) (CA INDEX NAME)

RN 135578-20-0 CAPLUS

CN 1H-Indole-3-acetic acid, α -(dicyanomethyl)-1-methyl-2-phenyl- α -(trifluoromethyl)-, ethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 38 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1991:471633 CAPLUS

DOCUMENT NUMBER:

115:71633

TITLE:

Preparation of pyrrolopyrimidines as antitumor agents

INVENTOR(S):

Akimoto, Hiroshi; Hitaka, Takenori

PATENT ASSIGNEE(S):

Takeda Chemical Industries, Ltd., Japan

SOURCE:

Eur. Pat. Appl., 27 pp.

DOCUMENT TYPE:

CODEN: EPXXDW

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 418924	, A2	19910327	EP 1990-118202	19900921 <

EP 418924 19911023 А3 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE JP 03173890 Α 19910729 JP 1990-249615 19900918 <--CA 2025830 19910322 Α1 CA 1990-2025830 19900920 <--US 5354754 Α 19941011 US 1993-46917 19930414 <--PRIORITY APPLN. INFO.: JP 1989-245998 19890921 Α US 1990-585950 B1 19900921 OTHER SOURCE(S): MARPAT 115:71633 GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Pyrrolopyrimidines [I; R1, R2 = H, ester residue; X = NH2, OH, SH; Y = H, OH; Z = (substituted) C2-4 divalent radical; Z1 = (substituted) divalent heterocycle residue, alkylene; dotted line indicates saturation or unsatn] are prepared Acetal II (1.32 g) (preparation given) was dissolved in CF3CO2H containing

H2O with stirring at room temperature to give quant. salt III, which was dissolved with di-Et glutamate HCl in DMF and the solution was treated with 0.514 g H2NP(O)(OEt)2 and Et3N in DMF at room temperature to give 1.11 g diester

IV (R1 = R2 = Et) (V). Saponification of 1.05 g V in THF gave 0.826 g acid IV (R1 $\,$

= R2 = H), which showed IC50 of 0.00043 $\mu g/mL$ against human epidermoid carcinoma KB cells.

IT 135110-11-1P 135111-93-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, in preparation of antitumor agent)

RN 135110-11-1 CAPLUS

CN Nonanedioic acid, 2-(dicyanomethyl)-, dimethyl ester (9CI) (CA INDEX NAME)

RN 135111-93-2 CAPLUS

CN 2-Thiophenepentanoic acid, α-(dicyanomethyl)-5-[(1,1-dimethylethoxy)carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 39 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1991:429914 CAPLUS

DOCUMENT NUMBER: 115:29914

TITLE: Preparation of N-[[(pyrrolopyrimidinylethyl)amino]benz

INVENTOR(S):

SOURCE:

oyl]glutamates and analogs as antitumor agents Akimoto, Hiroshi; Hitaka, Takenori; Miwa, Tetsuo

PATENT ASSIGNEE(S):

Takeda Chemical Industries, Ltd., Japan

Eur. Pat. Appl., 51 pp.

CODEN: EPXXDW

DOCUMENT TYPE: LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
EP 400562	A1	19901205	EP 1990-110131	19900529	<
EP 400562	B1	19960821	•		
R: AT, BE, CH,	DE, DK	, ES, FR, GB	, GR, IT, LI, LU, NL	, SE	
JP 04009382	Α	19920114	JP 1990-136345	19900525	<
JP 3015957	B2	20000306			
CA 2017604	A1	19901129	CA 1990-2017604	19900528	<
AT 141603	T	19960915	AT 1990-110131	19900529	<
PRIORITY APPLN. INFO.:			JP 1989-135642	A 19890529	
			JP 1989-246209	A 19890920	
			JP 1990-93370	A 19900409	
OTHER SOURCE(S):	MARPAT	115.29914			

The title compds. [I; R = (CR1R2)iZ1(CR4R5)jZ2CONHCH(CO2R6)CH2CH2CO2R7; R1, R2, R4, R5 = H, hydrocarbyl, bond; R6, R7 = H, alkyl, (un)substituted Ph, PhCH2; R8, R9 = H; R8R9 = bond; X = NH2, OH, SH; Y = H, OH; Z1 = O, SOn, (alkyl)imino, etc.; Z2 = (un)substituted alkylene, divalent cyclic group; i,j = 0-3 (i + j = 1-3); n = 0-2] were prepared Thus, 4-(EtO2C)C6H4NMeCH2CH2CH(CO2Me)CH(CN)2 (preparation given) was cyclocondensed with guanidine and the product reduced to give, as 1 of 2 products, anilinoethylpyrrolopyrimidine I [R = CH2CH2NMeC6H4(COR10)-4; R8R9 = bond, X = NH2, Y = H] (II; R10 = OEt) which was condensed with di-Et L-glutamate to give, after saponification, L-II [R10 = NHCH(CO2H)CH2CH2CO2H]. The latter

had

GI

IC50 of 0.0013 μM against human epidermoid carcinoma KB cell growth in vitro.

IT 133719-38-7P 133719-41-2P 133719-45-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, in preparation of antitumor agents)

RN 133719-38-7 CAPLUS

CN Benzoic acid, 4-[[4,4-dicyano-3-(methoxycarbonyl)butyl]methylamino]-, ethyl ester (9CI) (CA INDEX NAME)

RN 133719-41-2 CAPLUS
CN Benzoic acid, 4-[[4,4-dicyano-3-(ethoxycarbonyl)butyl][(1,1-dimethylethoxy)carbonyl]amino]-, ethyl ester (9CI) (CA INDEX NAME)

RN 133719-45-6 CAPLUS
CN Benzoic acid, 4-[4,4-dicyano-3-(ethoxycarbonyl)butoxy]-, ethyl ester (9CI)
(CA INDEX NAME)

L6 ANSWER 40 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1991:429194 CAPLUS

DOCUMENT NUMBER: 115:29194

TITLE: Synthesis of esters of 3,3-dicyano-2-

(trifluoromethyl)acrylic acid and their reactions with

aryl amines

AUTHOR(S): Tyutin, V. Y.; Chkanikov, N. D.; Kolomiets, A. F.;

Fokin, A. V.

CORPORATE SOURCE: Inst. Organoelem. Compd., Moscow, 117813, USSR

SOURCE: Journal of Fluorine Chemistry (1991), 51(3),

323-34

CODEN: JFLCAR; ISSN: 0022-1139

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 115:29194

GT

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Title acrylates (NC)2C:C(CF3)CO2R (I; R = Me, Et) were prepared by the condensation of CH2(CN)2 with CF3COCO2R in presence of ZnCl2. Reaction of I with aromatic amines was investigated. Thus, 2,6-dimethylaniline reacted with I in CHCl3 to give adduct II. 2,5-Dimethoxyaniline, and Ph2NH gave similar adducts. o- And m-C6H4(NH2)2 reacted with I to give cyclocondensation products, quinoxalinone III and indoline IV resp. 4-R1C6H4NHNH2 (R1 = H, NO2) gave pyrazolines V on cyclocondensation with I. Reaction of I with 3-aminopyrazole gave pyrazolopyridines VI.

IT 134641-38-6P 134641-39-7P 134641-40-0P 134641-41-1P 134641-42-2P 134641-43-3P

134641-44-4P

RN 134641-38-6 CAPLUS

CN Benzeneacetic acid, 4-amino- α -(dicyanomethyl)-3,5-dimethyl- α -(trifluoromethyl)-, methyl'ester (9CI) (CA INDEX NAME)

RN 134641-39-7 CAPLUS

CN Benzeneacetic acid, 4-amino- α -(dicyanomethyl)-3,5-dimethyl- α -(trifluoromethyl)-, ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & F3C & CN \\ & & | & | \\ & & C-CH-CN \\ \hline & & C-OEt \\ & & | \\ & & O \end{array}$$

RN 134641-40-0 CAPLUS

CN Benzeneacetic acid, 4-amino- α -(dicyanomethyl)-2,5-dimethoxy- α -(trifluoromethyl)-, methyl ester (9CI) (CA INDEX NAME)

RN 134641-41-1 CAPLUS

CN Benzeneacetic acid, 4-amino- α -(dicyanomethyl)-2,5-dimethoxy- α -(trifluoromethyl)-, ethyl ester (9CI) (CA INDEX NAME)

RN 134641-42-2 CAPLUS

CN Benzeneacetic acid, α -(dicyanomethyl)-4-(phenylamino)- α (trifluoromethyl)-, methyl ester (9CI) (CA INDEX NAME)

RN 134641-43-3 CAPLUS

CN Benzeneacetic acid, α -(dicyanomethyl)-4-(dimethylamino)- α -(trifluoromethyl)-, methyl ester (9CI) (CA INDEX NAME)

RN 134641-44-4 CAPLUS

CN Benzeneacetic acid, α -(dicyanomethyl)-4-(dimethylamino)- α -(trifluoromethyl)-, ethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 41 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1991:247080 CAPLUS

DOCUMENT NUMBER: 114:247080

TITLE: Reaction of α, β -unsaturated nitriles with

phosphorus ylides

AUTHOR(S): Abdou, Wafaa M.; Ganoub, Neven A. F.

CORPORATE SOURCE: Natl. Res. Cent., Dokki, Egypt

SOURCE: Chemistry & Industry (London, United Kingdom) (

1991), (6), 217-18

CODEN: CHINAG; ISSN: 0009-3068

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 114:247080

GI

HO
$$C(CN)_2$$
 $X=C$
 CR
 C

AB Reaction of unsatd. nitrile I with ROCC-HP+Ph3 (R = OMe, OEt, Ph) in benzene at 25° gave 75-80% addition products II (X = PPh3). On heating II (X = PPh3) to 200° they underwent an intramol. Wittig reaction to give arenopyrans III. Heating phosphorane II (R = OMe, X = PPh3) with BzH gave II (X = CHPh).

IT 133973-19-0P 133973-20-3P
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
 (Reactant or reagent)

(preparation and intramol. Wittig reaction of)

RN 133973-19-0 CAPLUS

CN 9-Phenanthrenepropanoic acid, β , β -dicyano-10-hydroxy- α - (triphenylphosphoranylidene)-, methyl ester (9CI) (CA INDEX NAME)

RN 133973-20-3 CAPLUS

CN 9-Phenanthrenepropanoic acid, β , β -dicyano-10-hydroxy- α - (triphenylphosphoranylidene)-, ethyl ester (9CI) (CA INDEX NAME)

IT 133973-25-8P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 133973-25-8 CAPLUS

CN 9-Phenanthrenepropanoic acid, β , β -dicyano-10-hydroxy- α - (phenylmethylene)-, methyl ester (9CI) (CA INDEX NAME)

ANSWER 42 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1991:121991 CAPLUS

DOCUMENT NUMBER: 114:121991

TITLE:

Reactions of malononitrile with acetylenic esters and

ketones [Erratum to document cited in

CA113(25):231170y]

AUTHOR(S):

Kandeel, Kamal A.; Vernon, John M.; Dransfield, Trevor

A.; Fouli, Fouli A.; Youssef, Ahmed S. A.

CORPORATE SOURCE:

Dep. Chem., Univ. York, Heslington/York, YO1 5DD, UK

SOURCE:

Journal of Chemical Research, Synopses (1990

), (12), 406

CODEN: JRPSDC; ISSN: 0308-2342

DOCUMENT TYPE:

Journal

LANGUAGE:

English

An error in the structure for compound 13 has been corrected. The error was not

reflected in the abstract or the index entries.

IT 130747-61-4P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of (Erratum))

RN 130747-61-4 CAPLUS

2-Butenedioic acid, 2-(dicyanomethyl)-, dimethyl ester (9CI) CN (CA INDEX

ANSWER 43 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1991:82446 CAPLUS

DOCUMENT NUMBER:

114:82446

TITLE:

Novel pyrrolo[2,3-d]pyrimidine antifolates: synthesis

and antitumor activities

AUTHOR(S):

Miwa, Tetsuo; Hitaka, Takenori; Akimoto, Hiroshi;

Nomura, Hiroaki

CORPORATE SOURCE:

Res. Dev. Div., Takeda Chem. Ind., Ltd., Osaka, 532,

Japan

SOURCE:

Journal of Medicinal Chemistry (1991),

34(2), 555-60

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 114:82446

GI

$$_{\text{H}_{2}\text{N}}^{\text{NH}_{2}}$$
 $_{\text{N}}^{\text{R}}$ $_{\text{CO}_{2}\text{CMe}_{3}}^{\text{R}}$ $_{\text{III}}^{\text{R}}$

AB Title compds. I (R = H, Me; R1 = H, Et; R2R3 = bond, R2 = R3 = H) were prepared as antifolates. A key step was the cyclocondensation of dicyano compound II (R = H, Me) with guanidine-HCl to give pyrrolo[2,3-d]pyrimidines III. III were prepared in several steps from p-RCOC6H4CO2CMe2 and CH3CH:CHCO2Et or BrCH2CH:CHCO2Et. These antifolates were more growth-inhibitory by about 1 order of magnitude than methotrexate (MTX) against KB human epidermoid carcinoma cells and A549 human nonsmall cell lung carcinoma cells in in vitro culture.

IT 125991-47-1P 130351-33-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and cyclocondensation of, with guanidine)

RN 125991-47-1 CAPLUS

CN Benzenepentanoic acid, α -(dicyanomethyl)-4-[(1,1-dimethylethoxy)carbonyl]- δ -methyl-, ethyl ester (9CI) (CA INDEX NAME)

RN 130351-33-6 CAPLUS

CN Benzenepentanoic acid, α -(dicyanomethyl)-4-[(1,1-dimethylethoxy)carbonyl]-, ethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 44 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1990:631170 CAPLUS

DOCUMENT NUMBER: 113:231170

TITLE: Reactions of malononitrile with acetylenic esters and

ketones

AUTHOR(S): Kandeel, Kamal A.; Vernon, John M.; Dransfield, Trevor

A.; Fouli, Fouli A.; Youssef, Ahmed S. A.

CORPORATE SOURCE: Dep. Chem., Univ. York, Heslington/York, YO1 5DD, UK

SOURCE: Journal of Chemical Research, Synopses (1990)

), (9), 276-7

CODEN: JRPSDC; ISSN: 0308-2342

DOCUMENT TYPE: Journal

LANGUAGE: English
OTHER SOURCE(S): CASREACT 113:231170

GI CASREAC

OMe CN

AB The addition of malononitrile to acetylenic esters and acetylenic ketones catalyzed by sodium alkoxides gave 3- and 5-cyano-2-pyridones, e.g., I, 2-cyano- and 2,6-dicyanoaniline, and other products.

IT 130747-61-4P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 130747-61-4 CAPLUS

CN 2-Butenedioic acid, 2-(dicyanomethyl)-, dimethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 45 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1990:497432 CAPLUS

DOCUMENT NUMBER: 113:97432 ·

TITLE: Quinolone antibacterial agents substituted at the

7-position with spiroamines. Synthesis and

structure-activity relationships

AUTHOR(S): Culbertson, Townley P.; Sanchez, Joseph P.; Gambino,

Laura; Sesnie, Josephine A.

CORPORATE SOURCE: Parke-Davis Pharm. Res. Div., Warner-Lambert Co., Ann

Arbor, MI, 48105, USA

SOURCE: Journal of Medicinal Chemistry (1990),

33(8), 2270-5

Journal

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE:

LANGUAGE: English

OTHER SOURCE(S): CASREACT 113:97432

GΙ

AB Fluoroquinolone antibacterials having the 7-position (10-position of pyridobenzoxazines) substituted with 2,7-diazaspiro[4.4]nonane, 1,7-diazaspiro[4.4]nonane, or 2,8-diazaspiro[5.5]undecane (e.g. I (X = CF, CH, N) were prepared and their biol. activities were compared with piperazine and pyrrolidine substituted analogs. Most exhibited potent Gram-pos. and Gram-neg. activity, especially when side chain was N-alkylated. Thus, the quinolinecarboxylic acid II was treated with 2-methyl-2,7-diazaspiro[4.4]nonane to give I (X = CH).

IT 77415-69-1

RL: RCT (Reactant); RACT (Reactant or reagent)
 (reductive cyclization of)

RN 77415-69-1 CAPLUS

CN Pentanedioic acid, 3,3-dicyano-, diethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 46 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1990:235871 CAPLUS

DOCUMENT NUMBER: 112:235871

TITLE: New gem-dicyanocyclobutane-containing hydroxyesters AUTHOR(S): Mori, Shojhi; Kakuchi, Toyoji; Padias, Anne Buyle;

Hall, H. K., Jr.

CORPORATE SOURCE: Chem. Dep., Univ. Arizona, Tucson, AZ, 85721, USA

SOURCE: Journal of Polymer Science, Part A: Polymer Chemistry

(1990), 28(3), 551-8

CODEN: JPACEC; ISSN: 0887-624X

DOCUMENT TOURS

DOCUMENT TYPE: Journal LANGUAGE: English

AB Six gem-dicyanocyclobutanes containing carbomethoxy and hydroxyl/acetoxy functions were synthesized by cycloaddn. of the appropriate vinyl ethers

or alkoxystyrenes to Me $\beta,\beta\text{-dicyanoacrylate}.$ They were too thermally liable to allow polycondensation to potentially piezoelec. linear polyesters.

IT 127396-28-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and attempted polymerization of)

RN 127396-28-5 CAPLUS

CN 1,3-Dioxolane-2-propanoic acid, α -(dicyanomethyl)-, methyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 47 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1990:158968 CAPLUS

DOCUMENT NUMBER:

112:158968

TITLE:

Preparation of N-[(pyrrolopyrimidinylalkyl)benzoyl]glu

tamates as neoplasm inhibitors

INVENTOR(S):

Akimoto, Hiroshi; Hitaka, Takenori; Miwa, Tetsuo

PATENT ASSIGNEE(S):

Takeda Chemical Industries, Ltd., Japan

SOURCE:

Eur. Pat. Appl., 24 pp.

DOCUMENT TYPE:

CODEN: EPXXDW Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	TENT NO.			KINI	DATE		PLICATION NO.		DATE	
EP	334636			A2	19890927		 1989-302851			<
EP	334636			A3	19910502				•	
ΕP	334636			В1	19961023					
					FR, GB,	GR, I	r, Lļ, Lu, NL,	SE		
ИО	8901206 169490	•		Α	19890925	ИО	1989-1206		19890320	<
ИО	169490		•	В	19920323		•	•		
ИО	169490 169490 4997838			С	19920701					
US	4997838			Α	19910305	US	1989-326901		19890321	<
DK	8901437			Α	19890925	DK	1989-1437		19890322	<
	173980			В1	20020325				•	
AΤ	144513				19961115	AT	1989-302851	•	19890322	<
ES	2092994			T3 C	19961216	ES	1989-302851		19890322	<
CA	1340794				19991019	CA	1989-594699		19890323	<
CN	1037513			Α	19891129	CN	1989-101681		19890324	<
CN	1029970			В	19951011					
HU	51624			A2	19900528	· HU	1989-1517		19890324	<
HU	203105			В	19910528					
.JP	02167281			Α	19900627	JP	1989-72235		19890324	<
JP	07005599			В	19950125					
HU	55396			A2	19910528	HU	1990-8458		19890324	<
HU	215928			В	19990329					
US	5106974				19920421	US	1990-578258		19900906	<
NO	9100661			A	19890925		1991-661		19910219	
NO	178304			В	19951120					
NO	178304			С	19960228					

US 5296600 US 5539113 PRIORITY APPLN. INFO.:		19960723 US JI JI NO	5 1992-824106 5 1993-161533 2 1988-71149 2 1988-245379 0 1989-1206 5 1989-326901	A A1	19920122 19931206 19880324 19880929 19890320	-
			3 1989-326901		19890321	
			5 ·1990-578258		19900906	
			5 1992-824106		19920122	
OMBED COUDCE/C/.	~~~~~~~~~~~	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	MADDAM 110.1500/0			

OTHER SOURCE(S):

CASREACT 112:158968; MARPAT 112:158968

AB The title compds. [I; R = H, F, alkyl, alkenyl, alkynyl; R3, R4 = H; R3R4
= bond; R5 = C6H4(CONHCHR1CH2CH2R2)-4; R1,R2 = (un)esterified CO2H; X =
NH2, OH; Y = H, NH2, OH; n = 2-4] were prepared Thus, 4(Me3CO2C)C6H4(CH2)3CH[CH(CN)2]CO2Me (preparation given) was refluxed 28 h with
(H2N)2C:NH.HCl in Me3COH containing Me3COK to give I [R = R3 = H, R5 =
C6H4(CO2CMe3)-4, X = NH2, n = 3] (II; R4Y = O) which was hydrogenated to
II (R4 = Y = H). The latter was hydrolyzed and the product condensed with
di-Et L-glutamate to give title compound III (R6 = Et, X = NH2) which was
hydrolyzed to III (R6 = H, X = OH) which had IC50 of 0.0006 μg/mL
against human nasopharyngeal cancer KB cells in vitro.

IT 125991-38-0P 125991-47-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, in preparation of neoplasm inhibitors)

RN 125991-38-0 CAPLUS

CN Benzenepentanoic acid, α -(dicyanomethyl)-4-[(1,1-dimethylethoxy)carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 125991-47-1 CAPLUS

CN Benzenepentanoic acid, α -(dicyanomethyl)-4-[(1,1-

dimethylethoxy)carbonyl]- δ -methyl-, ethyl ester (9CI) (CA INDEX NAME)

ANSWER 48 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1989:415389 CAPLUS

DOCUMENT NUMBER:

111:15389

TITLE:

Color photothermographic elements containing leuco

compounds

INVENTOR(S):

Sakizadeh, Kumars; Weigel, David C.; Grieve, Duncan;

Poon, Stephen S. C.; Thien, Tran V.

PATENT ASSIGNEE(S):

Minnesota Mining and Manufacturing Co., USA

SOURCE:

Eur. Pat. Appl., 35 pp. CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	TENT NO.		KIND		DATE	AP:	PLICATION NO.		DATE	
	294099	_	A2		19881207	EP	1988-304771		19880526	<
EP	294099		A3		19890531					•
EP	294099		B1		19930818					
•	R: BE, DE	FR,	GB,	ΙT						
US	4883747		Α		19891128	US	1988-200665		19880531	<
CA	1331107		С		19940802	CA	1988-568396		19880602	<
ŪΑ	8817345		Α		19881208	ΑU	1988-17345		19880603	<
AU	606162		B2		19910131					
JP	02032332		Α		19900202	JP	1988-145566		19880613	<
JP	2590204		B2		19970312					
US	4923792		Α		19900508	US	1989-368566		19890620	<
PRIORIT	Y APPLN. INE	·. o				GB	1987-12961	Α	19870603	
						US	1988-200665	A1	19880531	

OTHER SOURCE(S): CASREACT 111:15389; MARPAT 111:15389

A photothermog. material comprises Ag halide in reactive association with a Ag .salt of an organic acid and a color-generating reducing agent which is a leuco compound oxidizable by Ag ions into a colored dye of the formula ArR1C(:C(R5)C(R4):)nCR2R3 [n = 0-2; R1 = H, CN, C1-5 alkyl, aryl, CO2R6; R6 = C1-5 alkyl or aryl; R2, R3 = CN, NO2, CO2R6, SO2R6, COR6; R3 and R2may combine together to form a ring; R4, R5 = H, CN, C1-5 alkyl, or R4 and R5 together may form a ring; Ar = thienyl, furyl, phenyl]. The material produces images with improved color stability. Thus, a green-yellow image was produced with a photothermog. material incorporating leuco form of (p-dimethylaminobenzylidene)dimethylbarbituric acid.

IT 121246-61-5

RL: USES (Uses)

(photothermog. material containing, for improved image stability)

RN 121246-61-5 CAPLUS

ANSWER 49 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1988:570272 CAPLUS

DOCUMENT NUMBER:

109:170272

TITLE:

Synthesis and cognition-activating properties of some

mono- and bicyclic lactam derivatives

AUTHOR(S):

Altomare, Cosimo; Carotti, Angelo; Casini, Giovanni; Cellamare, Saverio; Ferappi, Marcello; Gavuzzo,

Enrico; Mazza, Fernando; Pantaleoni, Giancarlo;

Giorgi, Raffaele

CORPORATE SOURCE:

Dip. Farm.-Chim., Univ. Bari, Bari, Italy

SOURCE:

Journal of Medicinal Chemistry (1988),

31(11), 2153-8

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 109:170272

GI

AB Upon reductive cyclization cyano esters EtO2CCH2CH(CO2Et)CHRCN (R = CO2Et, cyano) and NCCH2CH(CN)CH(CO2Et)2 yielded piperidones and perhydropyrrolo[3,4-c]pyridine lactams I, II and III, resp. generally as a mixture of diastereomeric cis-trans forms. X-ray crystallog. anal. were carried out on cis-II and III. A series of neuropsychopharmacol. tests performed on I, II; and III indicated that they are generally nontoxic even at high doses (up to 1000 mg/kg i.p.)9. The cognition activating properties of lactams cis- and trans-I, cis-II, and III were evaluated in enhancing retention for passive avoidance learning in rats without and after electroconvulsive shock (ECS); compds. cis-I and III were found to be more potent than piracetam in the amnesia-reversal testing.

82584-86-9 IT

> RL: RCT (Reactant); RACT (Reactant or reagent) (reductive cyclization of, cyclic lactams from)

RN82584-86-9 CAPLUS

CN Butanedioic acid, (dicyanomethyl) -, diethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 50 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1988:94172 CAPLUS

DOCUMENT NUMBER: 108:94172

TITLE: Addition of ylidenemalononitriles onto dimethyl

acetylenedicarboxylate

AUTHOR(S): Gewald, Karl; Hain, Ute; Gruner, Margit

Sekt. Chem., Tech. Univ. Dresden, Dresden, DDR-8027, CORPORATE SOURCE:

Ger. Dem. Rep.

MeO₂C

Me

SOURCE: Zeitschrift fuer Chemie (1987), 27(1), 32-4

CODEN: ZECEAL; ISSN: 0044-2402

Ph

CN

NH₂

DOCUMENT TYPE: Journal German LANGUAGE:

OTHER SOURCE(S): CASREACT 108:94172

IV

GT

AΒ (NC)2C:CMeCH2CO2Me and MeO2CC.tplbond.CCO2Me (I) in the presence of K2CO3 cycloadded to give 49% aniline II. (NC) 2C: CPhMe and I, treated with Et3N, gave 30% dihydroquinoline III, which was aromatized by heating at 270° in No-MeOH to 80% quinolinecarboxylate IV. (NC)2C:CPhEt and I is a respectively gave 29% (NC) 2C:CPhCHMeCH(CO2Me) CH(CO2Me) C(CN) 2CPh:CHMe, which cyclized to cyclopentapyridinedicarboxylate V. IT

Ph

V

112754-03-7P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and thermal intramol. cyclization of)

RN 112754-03-7 CAPLUS

Butanedioic acid, 2-(3,3-dicyano-1-methyl-2-phenyl-2-propenyl)-3-(1,1-CN dicyano-2-phenyl-2-butenyl)-, dimethyl ester (9CI) (CA INDEX NAME)

CAPLUS COPYRIGHT 2007 ACS on STN ANSWER 51 OF 68

ACCESSION NUMBER:

1987:597249 CAPLUS

DOCUMENT NUMBER:

107:197249

TITLE:

Influence of the solvent on the nature of a

tetramethylene biradical intermediate

AUTHOR(S):

Padias, Anne Buyle; Hall, H. K., Jr.

CORPORATE SOURCE:

Chem. Dep., Univ. Arizona, Tucson, AZ, 85721, USA

SOURCE:

Journal of Organic Chemistry (1987), 52(20),

4536-9

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 107:197249

In the spontaneous thermal reactions of p-methoxystyrene and Me 3,3-dicyanoacrylate, several reaction products are observed: a 1/1 alternating copolymer, a double Diels-Alder adduct, and the cyclobutane adduct. In dipolar aprotic solvents, no polymerization occurs, and the double Diels-Alder adduct is favored; in protic polar solvents cyclobutane formation competes with copolymn. In nonpolar solvents, copolymn. dominates. A biradical tetramethylene species is proposed as the key intermediate. In polar solvents, this biradical exhibits considerable polar character, and Coulombic attraction between the termini favors the coiled or gauche conformation, leading preferentially to cycloadducts. nonpolar solvents, the trans conformation initiates the polymerization The

main

factors influencing the products are the solvent polarity and the ability of the solvent to interact with the biradical.

IT 110193-00-5

RL: PRP (Properties)

(conformation and spin and electron d. of, solvent effects on)

110193-00-5 CAPLUS RN

1,4-Butanediyl, 1,1-dicyano-2-(methoxycarbonyl)-4-(4-methoxyphenyl)- (9CI) CN (CA INDEX NAME)

IT 110193-05-0P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 110193-05-0 CAPLUS

Poly[1,1-dicyano-2-(methoxycarbonyl)-3-(4-methoxyphenyl)-1,4-butanediyl] CN

L6 ANSWER 52 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1987:196834 CAPLUS

DOCUMENT NUMBER:

106:196834

TITLE:

Cationic polymerization of nitrogen-containing

electron-rich vinyl monomers by electrophilic olefins

and their cyclobutane cycloadducts

AUTHOR(S):

Abdelkader, Mohamed; Padias, Anne Buyle; Hall, H. K.,

Jr.

CORPORATE SOURCE:

Chem. Dep., Univ. Arizona, Tucson, AZ, 85721, USA

SOURCE:

Macromolecules (1987), 20(5), 944-8 CODEN: MAMOBX; ISSN: 0024-9297

DOCUMENT TYPE:

Journal

LANGUAGE:

English

The major pathways for the reactions of very electron-rich N-containing olefins with several electrophilic olefins were studied. N-Ethyl-3-vinylcarbazole (I) [1486-07-3], N-vinylcarbazole (II) [1484-13-5], and p-(dimethylamino)styrene (III) [2039-80-7] underwent kinetic cyclobutane formation with an electrophilic olefin without a leaving group, Me β , β -dicyanoacrylate (IV) [82849-50-1], and one with a weak β -leaving group, tetracyanoethylene (V) [670-54-2]. The third electrophilic olefin, β , β -dicyanovinyl chloride (VI) [10472-09-0], had a strong β -leaving group and readily initiated the

of

donor olefin was used, IV, V, and VI all initiated cationic homopolymn. of I and II, while III only led to oligomers, as it did with conventional Broensted initiators. Cationic initiation by their own cyclobutane adducts was observed for the very electron-rich monomers I and II. Postcyanovinylation of the formed polymers by the electrophilic olefins occurred. Incorporation of a β -leaving group enhanced the initiating ability of the electrophilic olefins and N-carbazyl and N-ethyl-3-carbazyl were overall the most effective donor substituents favoring cationic homopolymn.

cationic polymerization of I and II and oligomerization of III. If an excess

IT 107540-79-4P

RL: SPN (Synthetic preparation); PREP.(Preparation)

(preparation of, from dimethylaniline and dicyanovinyl compound)

RN 10.7540-79-4 CAPLUS

CN Benzeneacetic acid, α -(dicyanomethyl)-4-(dimethylamino)-, methyl

ester (9CI) (CA INDEX NAME)

ANSWER 53 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1987:33439 CAPLUS

DOCUMENT NUMBER:

106:33439

TITLE:

Synthesis of novel symmetric diamino acids

AUTHOR(S):

Reddy, P. Anantha; Erickson, Bruce W.

CORPORATE SOURCE:

Rockefeller Univ., New York, NY, 10021, USA

SOURCE:

Pept.: Struct. Funct., Proc. Am. Pept. Symp., 9th (

1985), 453-6

CODEN: 54ZNAJ

DOCUMENT TYPE:

Conference

LANGUAGE:

English

Sym diamino acids (H2NCH2)2CHCH2CO2H (Aab) 3,5-(H2NCH2X)2C6H3CO2H [X = null, CH2 (Bab)] were prepared from (NC)2CHCH2CO2CMe3 and

3,5-(BrCH2)2C6H3CO2Me. The N,N-bis(tert-butoxycarbonyl) derivative of Aab couples efficiently during solid-phase peptide synthesis. The

corresponding derivative of Bab is used in the synthesis of the protein

betabellin.

IT 105995-37-7P 105995-39-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

(preparation and hydrogenation of)

RN 105995-37-7 CAPLUS

CN Propanoic acid, 3,3-dicyano-, 1,1-dimethylethyl ester (9CI) (CA INDEX

RN 105995-39-9 CAPLUS

Propanoic acid, 3,3-dicyano-, phenylmethyl ester (9CI) (CA INDEX NAME)

CAPLUS COPYRIGHT 2007 ACS on STN ANSWER 54 OF 68

ACCESSION NUMBER:

1986:496793 CAPLUS

DOCUMENT NUMBER:

105:96793

TITLE:

Zwitterionic tetramethylenes as the common.

intermediates in the cycloaddition and polymerization

reactions of N-vinylcarbazole with electrophilic tetrasubstituted ethylenes: a new explanation for

charge-transfer initiation

AUTHOR(S):

Gotoh, Tetsuya; Padias, Anne Buyle; Hall, H. K., Jr.

CORPORATE SOURCE:

Chem. Dep., Univ. Arizona, Tucson, AZ, 85721, USA

Journal of the American Chemical Society (1986

), 108(16), 4920-31

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: LANGUAGE:

SOURCE:

Journal English

OTHER SOURCE(S):

CASREACT 105:96793

AB The reactions of N-vinylcarbazole (I) with electrophilic tetrasubstituted ethylenes were examples of reactions whose outcomes are manipulated by changes in concentration, structure, and working procedure to form either small mols. (cyclobutanes, 1-butenes) or poly(vinylcarbazole). Equivalent concns. and evaporative workup (organic chemists' conditions) lead to small mols.; a large excess of I and precipitative workup give polymer. The mechanism involves gauche and trans zwitterionic tetramethylenes as intermediates. The former gives cyclobutane reversibly. The latter gives 1-butenes intramol. or adds monomers to form cyclohexanes or eventually polymer. The organic chemical and polymer chemical are unified on this basis. Extensive stereochem. and kinetic support for these propositions is given. Two other proposed mechanisms for these charge-transfer initiations are excluded.

RN 96735-90-9 CAPLUS

CN Propanedioic acid, [2-(9H-carbazol-9-yl)ethenyl](dicyanomethyl)-, dimethyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 102852-12-0 CAPLUS

CN 9H-Carbazole-9-butanoic acid, α -cyano- α -(dicyanomethyl)- γ -methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 102852-13-1 CAPLUS

CN 9H-Carbazole-9-pentanoic acid, α, β, β -tricyano- δ -methoxy-, methyl ester (9CI) (CA INDEX NAME)

RN 102852-14-2 CAPLUS

CN Propanedioic acid, [2-(9H-carbazol-9-yl)-2-methoxyethyl](dicyanomethyl)-, dimethyl ester (9CI) (CA INDEX NAME)

RN 102852-38-0 CAPLUS

CN 3-Butenoic acid, 4-(9H-carbazol-9-yl)-2-cyano-2-(dicyanomethyl)-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 102852-39-1 CAPLUS

CN 4-Pentenoic acid, 5-(9H-carbazol-9-yl)-2,3,3-tricyano-, methyl ester, (E)-(9CI) (CA INDEX NAME)

Double bond geometry as shown.

L6 ANSWER 55 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1985:406747 CAPLUS

DOCUMENT NUMBER: 103:6747

TITLE: Zwitterionic tetramethylene intermediates: a new

interpretation for "charge-transfer" initiation

AUTHOR(S): Hall, H. K., Jr.; Gotoh, T.

CORPORATE SOURCE: Dep. Chem., Univ. Arizona, Tucson, AZ, 85721, USA

SOURCE: Polymer Preprints (American Chemical Society, Division

of Polymer Chemistry) (1985), 26(1), 34-5

CODEN: ACPPAY; ISSN: 0032-3934

DOCUMENT TYPE: Journal LANGUAGE: English

AB Investigation of the initiation mechanism in polymerization of N-vinylcarbazole (I) [1484-13-5] in the presence of tetracyanoethylene [670-54-2] or di-Me 2,2-dicyanoethylene-1,1-dicarboxylate [82849-49-8] showed that neither the I-cyano compound charge transfer complexes nor the ion-radical pairs formed from them initiated polymerization. The initiating species was the gauche or trans tetramethylene zwitterion formed as an intermediate from the charge-transfer complex. This finding indicated that cyclobutanes initiated vinyl polymerization. The mechanism and the kinetics of the zwitterionic initiation were discussed.

IT 96735-90-9

RL: CAT (Catalyst use); USES (Uses)

(catalysts, for vinylcarbazole polymerization)

RN 96735-90-9 CAPLUS

CN Propanedioic acid, [2-(9H-carbazol-9-yl)ethenyl](dicyanomethyl)-, dimethyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

ANSWER 56 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

1983:488082 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 99:88082

Tetraoxo derivatives of perhydropyrrolo[3,4-c]pyridine TITLE: AUTHOR(S):

Ferappi, M.; Carotti, A.; Casini, G.; De Laurentis,

N.; Giardina, D.; Cingolani, G. M.; Gavuzzo, E.;

Mazza, F.

Ist. Chim. Farm. Tossicol., Univ. Bari, Bari, 70126, CORPORATE SOURCE:

Italy

SOURCE: Journal of Heterocyclic Chemistry (1983).,

20(2), 439-46

CODEN: JHTCAD; ISSN: 0022-152X

DOCUMENT TYPE: Journal

English LANGUAGE:

CASREACT 99:88082 OTHER SOURCE(S):

GI

CO2Et CONH₂ CO2Et CN EtO2C CN Ι II III

AΒ Michael adducts from di-Et furmarate with malonic esters or nitriles were cyclized to succinimide intermediates which, after glutarimide ring closure, afforded several N-Me and N-benzyl derivs. of cis-1,3,4,6-tetraoxoperhydropyrrolo[3,4-c]pyridine whose configuration was demonstrated by x-ray crystal structure anal. Thus, treating the adduct I with H2SO4 gave succinimide II which was treated with NaOEt in EtOH or

tosyl acid in xylene to give pyrrolopyridine III. 82584-86-9P IT

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and cyclization of, pyrrolidine from)

RN 82584-86-9 CAPLUS

CN Butanedioic acid, (dicyanomethyl)-, diethyl ester (9CI) (CA INDEX NAME)

0 OFt NC-CH-CH-CH2-C-OEt

ANSWER 57 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1982:582893 CAPLUS

DOCUMENT NUMBER: 97:182893

TITLE: Dimethyl 1,1-dicyanoethene-2,2-dicarboxylate, a new

electrophilic olefin

AUTHOR(S): Hall, H. K., Jr.; Sentman, R. C.

Dep. Chem., Univ. Arizona, Tucson, AZ, 85721, USA CORPORATE SOURCE:

SOURCE: Journal of Organic Chemistry (1982), 47(23),

4572-7

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE:

Journal

LANGUAGE:

English

dimethyl 1,1-dicyanoethene-2,2-dicarboxylate (I) [82849-49-8] was synthesized via a Knoevenagel condensation. I spontaneously copolymerizes with electron-rich olefins such as styrene [100-42-5] and p-methylstyrene [622-97-9]. In the copolymn., the bulky growing styryl radicals add to the dicyano-bearing carbon of I. Cyclobutane adducts are obtained in thermal reactions with styrene, p-methylstyrene, p-methoxystyrene [637-69-4], and vinyl ethers via a tetramethylene intermediate. formation occurs at the diester end of I due to the greater stabilization provided by the dicyano group and the minimal steric requirements of the attacking methylene.

IT 82849-58-9P

> RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and NMR spectra of)

RN 82849-58-9 CAPLUS

Benzenepropanoic acid, α -(1,1-dicyanoethyl)- β -ethyl-, methyl CN ester, polymer with ethenylbenzene (9CI) (CA INDEX NAME)

CM 1

CRN 82917-40-6 CMF C16 H18 N2 O2

CM 2

CRN 100-42-5 C8 H8 CMF

 $H_2C = CH - Ph$

ANSWER 58 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

1982:472220 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

97:72220

TITLE:

Contribution to the synthesis of the glutarimides.

AUTHOR(S):

Victory, Pedro; Jover, Jose Maria; Sempere, Julian Dep. Quim. Org., Inst. Quim. Sarria, Barcelona, Spain

CORPORATE SOURCE: SOURCE:

Afinidad (1981), 38(376), 491-5 CODEN: AFINAE; ISSN: 0001-9704

DOCUMENT TYPE:

Journal

LANGUAGE:

Spanish

OTHER SOURCE(S):

CASREACT 97:72220

GI

AB Glutarimides I [R = R1 = H, R2 = CO2Et, Ph, 3-furyl, 2-thienyl, Me; R = cyano, R1R2 = (CH2)5; R = Me, R1 = R2 = H] were prepared by treating CH2(CN)2 with R1R2C:CRCO2Et with or without isolation of (NC)2CHCR1R2CHRCO2Et, and acid hydrolysis of the enol ethers. Alternatively R1R2C:CRCO2Et was cyclized with NCCH2CONH2.

IT 82584-86-9P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and cyclization of)

RN 82584-86-9 CAPLUS

Butanedioic acid, (dicyanomethyl) -, diethyl ester (9CI) CN (CA INDEX NAME)

L6 ANSWER 59 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1981:442180 CAPLUS

DOCUMENT NUMBER:

95:42180

TITLE:

Absolute configuration of 2,7-diazaspiro[4,4] nonane.

A reassignment

AUTHOR(S):

Overberger, C. G.; Wang, David Wei; Hill, Richard K.;

Krow, Grant R.; Ladner, David W.

CORPORATE SOURCE:

Macromol. Res. Cent., Univ. Michigan, Ann Arbor, MI,

48109, USA

SOURCE:

Journal of Organic Chemistry (1981), 46(13),

2757-64

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 95:42180

GI

The absolute configuration of the axially dissym. spirane 2,7-AB diazaspiro[4,4] nonane (I), was elucidated as (R)-(-), (S)-(+) in CHCl3 by synthesis of both enantiomers from the centrodissym. intermediate II; the configuration of (R)-(-)-II was correlated with that of (S)-HO2CCMeEtCH2CO2H through the substituted pyrrolidine III. configuration thus established for the sulfonamide derivative IV is opposite to that derived earlier (Krow, G. and Hill, R. K., 1968). The source of the original error lies in the preparation of spiroimide V, which is accompanied by almost total racemization when carried out at high temps. A more direct, efficient synthesis of I is described, followed by resolution with dinitrodiphenic acid to give the optically pure enantiomers. Lowe's rule predicts correctly the absolute configurations of several I derivs. but not that of I itself.

 $\mathbf{T}\mathbf{I}$ 77415-69-1P

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and diazaspirononane derivative from)

RN 77415-69-1 CAPLUS ·

CN Pentanedioic acid, 3,3-dicyano-, diethyl ester (9CI) (CA INDEX NAME)

ANSWER 60 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

1978:50444 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

SOURCE:

88:50444

TITLE:

The chemistry of 2-oxopropanedinitrile (carbonyl

cyanide); XIX. The ene synthesis using

2-oxopropanedinitrile and 1,3-dicarbonyl compounds

AUTHOR(S):

Kociolek, K.; Leplawy, M. T.

CORPORATE SOURCE:

Inst. Org. Chem., Tech. Univ. Lodz, Lodz, Pol.

Synthesis (1977), (11), 778-80

CODEN: SYNTBF; ISSN: 0039-7881

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 88:50444

AΒ Reaction of CO(CN) 2 with RCOCH2COR1 (I; R = R1 = Ph, 2,4,6-Cl3C6H2, Me; R = Me, F3C, R1 = Ph) in ether at 0° was complete in 1 h and gave RCOCH(COR1)C(CN)2OH (II; R and R1 as before) in 100% yield. Reaction of

CO(CN)2 with I (R = Rl = OEt) at room temperature required 20 days and gave II

in 43-66% yield.

IT 65305-78-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, with aniline)

RN 65305-78-4 CAPLUS

CN Propanedioic acid, (dicyanohydroxymethyl)-, diethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 61 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1973:5

1973:545978 CAPLUS

DOCUMENT NUMBER:

79:145978

TITLE: INVENTOR(S):

O,O-Dialkylthiophosphoric acid pseudochalcogen acyls Koehler, Helmut; Gerats, Irmtraut; Eichler, Gerhard;

Kochmann, Werner

SOURCE:

Ger. (East), 14 pp.

CODEN: GEXXA8

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

gave
=
re
=

IT 50605-40-8P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 50605-40-8 CAPLUS

CN Propanoic acid, 3,3-dicyano-3-(dimethoxyphosphinothioyl)-, methyl ester (9CI) (CA INDEX NAME)

ANSWER 62 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1973:545936 CAPLUS

DOCUMENT NUMBER: 79:145936

TITLE: Reaction of some fluoroolefins with sodium cyanide AUTHOR(S): Dyatkin, B. L.; Sterlin, S. R.; Zhuravkova, L. G.;

Martynov, B. I.; Knunyants, I. L.

CORPORATE SOURCE: Inst. Elementoorg. Soedin., Moscow, USSR SOURCE: Zhurnal Organicheskoi Khimii (1973), 9(9),

1786-90

CODEN: ZORKAE; ISSN: 0514-7492

DOCUMENT TYPE: Journal . LANGUAGE: Russian

For diagram(s), see printed CA Issue. GI

(CF3)2C:CF2 reacted with 1 equiv of NaCN at -5 to 0° in dioxane AB containing H2O to give 38% (CF3)2C:CFCN (I), and with excess NaCN in THF containing H2O to give 55% NCC(CF3)2CH(CN)2, a C-H acid of pKa 2.12; analogous treatment of (CF3)2C:CFPh and (CF3)2CHCO2Et yielded 49% NCC(CF3)2CHPhCN and 51% EtO2CC(CN)(CF3)CH(CN)2, resp., after neutralization. Under similar conditions, CF3CF:CF2 afforded 59% CF3[C(CN)2]2Na, although its acid could not be isolated, and (CF3)2C:CFOEt gave 3% (CF3)2C:C(CN)OEt. reacted with H2SO4 and EtOH to give 25% (CF3)2C:CFCO2Et, with HCl in EtOH to give 20% HOC(CF3)2CHFCONH2, with Et2NH to give 43% (CF3)2C:C(CN)NEt2, with PhNH2 to give 60% (CF3)2CHC(CN):NPh, and with concentrated H2SO4 to give 84% iminolactone (II; R = H), which was converted to its Hg salt (II; R =1/2 Hg) with HgO in refluxing aqueous Me2CO.

IT 50616-04-1P

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 50616-04-1 CAPLUS

CN Propanoic acid, 2-cyano-2-(dicyanomethyl)-3,3,3-trifluoro-, ethyl ester (CA INDEX NAME)

ANSWER 63 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1965:462918 CAPLUS

DOCUMENT NUMBER: 63:62918 ORIGINAL REFERENCE NO.: 63:11492h

TITLE: Reaction of acetylenic esters with cyanoacetic ester

and pyridine

AUTHOR(S): Bamfield, P.; Crabtree, A.; Johnson, A. W.

CORPORATE SOURCE: Univ. Nottingham, UK

SOURCE: Journal of the Chemical Society (1965)

4355-62

CODEN: JCSOA9; ISSN: 0368-1769

DOCUMENT TYPE: Journal LANGUAGE: English

Modified structures are suggested for the yellow and the blue adducts from dimethyl acetylenedicarboxylate, Et cyanoacetate, and pyridine, which were originally prepared and formulated by Diels. The reaction of Me phenylpropiolate, Et cyanoacetate, and pyridine leads to a 1:1:1-adduct in which the pyridine has suffered ring-fission. Various reactions of the adducts are discussed.

1289-25-4 IT

(Derived from data in the 7th Collective Formula Index (1962-1966))

RN 1289-25-4 CAPLUS

CN 1,2,3,4-Pentanetetracarboxylic acid, 5,5-dicyano-, tetramethyl ester, compd. with pyridine (1:1) (8CI) (CA INDEX NAME)

CM 1

CRN 45287-28-3 CMF C15 H18 N2 O8

CM 2

CRN 110-86-1 CMF C5 H5 N



L6 ANSWER 64 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1964:484793 CAPLUS

DOCUMENT NUMBER: 61:84793

ORIGINAL REFERENCE NO.: 61:14826g-h,14827a-c

TITLE: 1-Halo-1,2,3,3-tetra(negatively substituted)propanes

and their salts

INVENTOR(S): Martin, Elmore L.

PATENT ASSIGNEE(S): E. I. du Pont de Nemours & Co.

SOURCE: 6 pp.
DOCUMENT TYPE: Patent
LANGUAGE: Unavailable

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE ·	APPLICATION NO.	DATE
,				
US 3133084		19640512	US	19600624 <
PRIORITY APPLN. INFO.:			US	19600624

AB Compds. of the general formula [XC(Z):C(Y)C(A)R]- M+ (I), where A, R, Y, Z are electron withdrawing groups such as CN, CO2Et, Bz, or SO2Ph, X is Cl or F, and M is H, Na, K, or a substituted ammonium ion, are dyes for natural and synthetic fibers. Thus, H2C(CN)2 79 in tetrahydrofuran (II) 220 was added with stirring to a dispersion 52 of 51.2% NaH in mineral oil and II 660 at 5-10° during 15 min., the mixture stirred 30 min., then dichlorofumaronitrile 44 in II 220 added during 15 min., II vaccum-distilled at 35-40°, the residual yellow solid dissolved in H2O 250, the pH adjusted to 8 with CO2, then Et4NBr 100 in H2O 200 parts added slowly with

stirring, the mixture cooled to 5°, and the yellow crystals of I (A = R = Y = X = CN, Z = Cl, M = Et4N) (III) filtered, washed with 1% Et4NBr, and then H2O. The cake was dissolved in H2O 3500 at 100°, decolorizing carbon 10 added, the solution clarified, cooled to 5°, the long yellow needles filtered, washed with H2O and air-dried, giving 70 parts III, m. 129-31°, λ maximum 387 m μ , ϵ = 18,200 (MeOH) yellow on cellulose acetate and nylon, brownish yellow on wool and silk. Similarly, other I were prepared as tabulated below: X, Z, Y, A, R, M, % yield, m.p., color, λ (m μ) maximum, ϵ ; Cl, PhN(CO-)2, CN, CN, Me4N, 31 230-5° (decompose), orange, 468, 12,200; Cl, CO2Me, CO2Me, CN, CN, Et4N, 82, 88-90°, yellow, 335, 29,400; Cl, CN, CN, CO2Et, CO2Et, H, 100, bl, 115-20°, yellow (Na salt), -, -; Cl, Bz, Bz, CN, CN, Me4N, 39, 210-12° (decompose), yellow, 416, 27,000; F, CF3, CF3, CN, CN, Pr4N, 81, 84-6°, yellow, -, -; Cl, CN, CN, CN, CN, Me4N, -, 217-18° (decompose), yellow, 386, 17,600; Cl, CN, CN, CN, CN, Pr4N, -, $74-6^{\circ}$ (decompose), yellow, 386, 18,100; Cl, CN, CN, CN, CN, Et3NH, -, $63-5^{\circ}$ (decompose), yellow, 387, 17,200; Cl, CN, CN, CN, CO2Et, Et4N, 56, 70-2°, yellow, 400, 15,700; Cl, CN, CN, CN, SO2C6H4Me-4, Me4N, 73, 124-6° (decompose), yellow, 387, 17,000; Cl, CN, CN, CN, Bz, Me4N, -, 159-61°, yellow, 414, 17,100; Cl, CN, CN, CN, Bz, Et4N, 30 118-19°, yellow, 420, 16,200; Cl, CN, CN, CN, CN, Pr4N, -, 109-10°, yellow, 412, 17,600; Cl, CF3, CF3, CN, CN, Et4N, 64, 84-5°, yellow, -, -; F, -CF2CF2-, CN, CN, Na, -, -, orange, -, -; Cl, CN, CN, Bz, Bz, Me4n, -, 167°9°, yellow, 422, 8000; Cl, CN, CN, CN, CONHPh, K, -, -, red, -, -; Cl, CN, CN, SO2Ph, SO2Ph, Me4N, -, -, yellow, -, -; Cl, CN, CN, Bz, CO2Et, H, 20, 97-8°, colorless, -, -; Cl, CN, CN, Bz, CO,2Et, Na, yellow; 98469-37-5P, Ammonium, tetraethyl, 1,2-dicarboxy-1-chloro-3,3-dicarboxy-1-chlo dicyanopropenide, dimethyl ester

IT RL: PREP (Preparation)

(preparation of)

RN98469-37-5 CAPLUS

CNTetraethylammonium 1,2-dicarboxy-1-chloro-3,3-dicyanopropenide, dimethyl ester (7CI) (CA INDEX NAME)

CM 1

CRN 98469-36-4 CMF C9 H6 C1 N2 O4

2 CM

CRN 66-40-0 C8 H20 N CMF

ANSWER 65 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1962:429396 CAPLUS

DOCUMENT NUMBER: 57:29396

ORIGINAL REFERENCE NO.: 57:5809h-i,5810c

TITLE:

Nitration of cyclohexanecarboxylic acid to caprolactam AUTHOR(S): Bigot, J. A.; Meijerink, Th. A. J.; Revallier, L. J.

CORPORATE SOURCE: Central Lab., Staatsmijnen, Geleen, Neth. SOURCE:

Recueil des Travaux Chimiques des Pays-Bas (

1962), 81, 363-4

CODEN: RTCPA3; ISSN: 0165-0513

DOCUMENT TYPE: Journal LANGUAGE: English

Cyclohexanecarboxylic acid (I) with nitryl hydropyrosulfate in oleum did not give the expected nitrocyclohexane, but 70% caprolactam (II) and a mixture of m-dinitrobenzene and nitrobenzene (total yield 22%, based on I). The mechanism of the reaction is unknown, but there is some evidence that removal of H2O from a nitro derivative is 1 of the steps involved. 1-Methyl-1-nitrocyclohexane with oleum gave a compound, C7H11NO (b2 79°, m. 48°), probably 1-methyl-1-nitrocyclohexene (or its rimer), a compound that could be isolated as such, since it could neither dehydrogenate to a C6H6 derivative, nor disproportionate and subsequently rearrange to II.

94211-18-4P, Acrylic acid, 3,3-dicyano-2-(dicyanomethyl)-, ethyl ΙT ester, compound with quinoline 94467-89-7P, Acrylic acid, 3,3-dicyano-2-(dicyanomethyl)-, ethyl ester, compound with NH3 RL: PREP (Preparation)

(preparation of)

RN 94211-18-4 CAPLUS

Acrylic acid, 3,3-dicyano-2-(dicyanomethyl)-, ethyl ester, compd. with CN quinoline (7CI) (CA INDEX NAME)

1 CM

CRN 94211-17-3 CMF C10 H6 N4 O2

CM 2

CRN 91-22-5 CMF C9 H7 N

RN 94467-89-7 CAPLUS

2-Propenoic acid, 3,3-dicyano-2-(dicyanomethyl)-, ethyl ester, ion(1-), CN

NH4 +

L6 ANSWER 66 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1962:429395 CAPLUS

DOCUMENT NUMBER: 57:29395
ORIGINAL REFERENCE NO.: 57:5809e-h

TITLE: Base-catalyzed ring opening of diethyl

1,1,2,2-tetracyanocyclopropane-3,3-dicarboxylate

AUTHOR(S): Regan, T. H.

CORPORATE SOURCE: E. I. du Pont de Nemours & Co., Wilmington, DE

SOURCE: Journal of Organic Chemistry (1962), 27,

2236-7

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

AB An example of a cyclopropane ring cleavage under very mild conditions was reported. CH2(CN)2 (6.6 g.) in 17.4 g. di-Et oxomalonate was left 3 hrs. with one drop of base catalyst; the solid was collected and shown to be di-Et dihydroxymalonate. Fractionation of the yellow oil gave 11.7 g. di-Et 1,1-dicyanoethylene-2,2-dicarboxylate, b1 86°, n24D 1.4628. An equimolar mixture of this compound and anthracene after heating at 150° gave crystalline product, m. 153.6-5.2° (alc.-H20). The product resulting from 38 g. CH2(CN)2 and 100 g. di-Et oxomalonate in 250 ml. alc. treated in the cold with 52 g. Br, the solution poured onto 1 kg. ice, and the oil crystallized when left overnight gave 71.5 g. di-Et 1,1,2,2-tetracyanocyclopropane-3,3-dicarboxylate (I), m. 129.6-31.2° (alc.-H2O). I (15 g.) was suspended in 500 ml. Et2O, treated with 15 g. dry NH3, stirred overnight and the mixture filtered to give 11.4 g. solid, m. 192-201° (decomposition). The filtrate evaporated and the residue stirred with CHCl3 gave 0.5 g. yellow powder, m. 203° (decomposition). The CHCl3 solution evaporated gave Et carbamate, m. 46.6-8.6°. The yellow powder was ammonium 1,1,3,3-tetracyano-2carbethoxypropenide (II). II in H2O treated with a concentrated aqueous solution of

quinclinium chloride gave quinolinium 1,1,3,3-tetracyano-2carbethoxypropenide, m. 111.5-12.5°. Recrystn. from H2O gave a

hydrate, m. 51-2°.

IT 94211-18-4P, Quinoline, compound with Et 3,3-dicyano-2-(dicyanomethyl)-acrylate 94467-89-7P, Acrylic acid, 3,3-dicyano-2-(dicyanomethyl)-, ethyl ester, compound with NH3 RL: PREP (Preparation)

(preparation of)

RN 94211-18-4 CAPLUS

CN Acrylic acid, 3,3-dicyano-2-(dicyanomethyl)-, ethyl ester, compd. with quinoline (7CI) (CA INDEX NAME)

CM 1

CRN 94211-17-3 CMF C10 H6 N4 O2

CM 2

CRN 91-22-5 CMF C9 H7 N

RN 94467-89-7 CAPLUS

CN 2-Propenoic acid, 3,3-dicyano-2-(dicyanomethyl)-, ethyl ester, ion(1-), ammonium (9CI) (CA INDEX NAME)

● NH4+

L6 ANSWER 67 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1940:18276 CAPLUS

DOCUMENT NUMBER:

34:18276

ORIGINAL REFERENCE NO.:

34:2801g-i,2802a-h

TITLE:

Synthesis of α,α -dimethyltricarballylic and 1-carboxy-cyclopentane-1- α -succinic and 1-carboxy-3-methylcyclopentane-1- α -succinic

acids

AUTHOR(S):

Desai, R. D.; Sahariya, G. S.

SOURCE:

Journal of the University of Bombay, Science:

Physical Sciences, Mathematics, Biological Sciences

and Medicine (1939), 8(Pt. 3), 235-8

CODEN: JUBSAS; ISSN: 0368-4644

DOCUMENT TYPE:

Journal Unavailable

LANGUAGE:

AB When in RR'C(CN)C(CN)CO2Et, R and R' together are cyclopentane or

methylcyclopentane rings, there are obtained with CH2BrCO2Et (I) excellent yields of tricarballylic acids which are characterized by their toluidide N-tolylimides. A mixture of the Et sodiocyanoacetate (II) from 24 g. cyanoacetate and 21 g. cyclopentanone cyanohydrin is allowed to stand for 48 h. After addition of 32 g. I the mixture is kept at room temperature for 2

and then refluxed until it is neutral. The EtOH is distilled off, the residue diluted with H2O and the oil extracted with ether, dried and distilled into

3 fractions, b4 90-120°, 120-65° and 185-7°. The 2nd fraction is retreated with I. Et 1-cyanocyclopentane-1-α-cyanosuccinate (III), b4 185-7°, is obtained in 45% yield. Hydrolysis of III with concentrated H2SO4 gives 1-carboxycyclopentanesuccinic acid (IV), m. 165° (cf. Chatterji, C. A. 31, 7409.7, found 159°). Its anilide N-phenylimide, prepared by heating IV with PhNH2 at 170-5° for 3 h., m. 156°; p-toluidide N-p-tolylimide m. 189-90°. Et 1-cyano-3-methylcyclopentane-1-α-cyanosuccinate (V), prepared as II, b12 205°. Hydrolysis of V gives 1-carboxy-3-methylcyclopentanesuccinic acid, m. 144°; its p-toluidide N-tolylimide m. 167°. Di-Et 2-methyl-3,3-dicyanobutane-3,4-dicarboxylate (VI), prepared from II, Me2C(OH)CN and I, in 45% yield, b5 176-8°. Saponification of VI with H2SO4 gives α,α-dimethyltricarballylic acid, m. 160° (C. found 156°). Its anilide N-phenylimide m. 140°; the p-toluidide N-p-tolylimide m.

IT 858794-64-6P, Glutaric acid, β , β -dicyano- α , α -dimethyl-, diethyl ester RL: PREP (Preparation) (preparation of)

RN 858794-64-6 CAPLUS

days

CN Glutaric acid, β , β -dicyano- α , α -dimethyl-, diethyl ester (4CI) (CA INDEX NAME)

L6 ANSWER 68 OF 68 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1938:911 CAPLUS

DOCUMENT NUMBER: 32:911

ORIGINAL REFERENCE NO.: 32:156d-i,157a-i,158a-e

TITLE: 2,3,-Dioxopyrrolines, mononuclear substances related

to isatin

AUTHOR(S): Mumm, Otto; Hornhardt, Hans

SOURCE: Berichte der Deutschen Chemischen Gesellschaft

[Abteilung] B: Abhandlungen (1937), 70E, ...

1930-47

CODEN: BDCBAD; ISSN: 0365-9488

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

AB 5-Phenyl-2,3-dioxopyrroline (I) (C. A. 5, 703) is so extraordinarily similar in appearance and chemical properties to isatin that it may be considered as a mononuclear isatin, and it was hoped that by means of this very reactive substance light might be obtained on some of the controversial questions on isatin, especially the structure of its salts and derivs. The earlier work on I was accordingly resumed and attention was directed to the preparation of analogs of I containing an aliphatic residue instead of

In order not to weaken the ring unnecessarily, a residue (hexyl) of rather high mol. weight was chosen. As a tertiary residue might also favor the stability of the ring, Me3C was selected for a 2nd series of expts. It was intended to prepare the new compds. by the earlier method. Pinacolone and C6H13COMe were condensed with HCO2R to the hydroxymethylene compds. which with NH2OH yielded the oxazoles through the intermediate oximes. The conversion of the oxazoles into the open-chain methylimide nitriles proceeded as expected and the formation from the nitriles of the desired pyrrolines with alc. HCl undoubtedly occurred, as evidenced by the appearance of the characteristic dark red color, but the products did not crystallize. The planned investigation was therefore continued with aromatic derivs., using p-tolyl instead of Ph compds. These tolyl compds., having higher m. ps., were considerably more stable and crystallized better. starting point was α -p-tolylisoxazole (II). That the nitrile obtained from II was really p-toluylpyruvonitrile methylimide, RC(OH):CHC(:NMe)CN(R = p-MeC6H4) (III), was shown by the reaction with MeMgI, which gave the normal product, RC(OH)MeCH2C(:NMe)C(:NMgI) (IV), and also the compound RC(NH2)MeCH2C(:NMe)C(:NMgI)Me(V) when the Grignard compound was decomposed with NH4Cl instead of water. RC(OH)MeCH2C(:NMe)C(OH)(NHMgI)M e (VI) was also formed by addition of H2O to IV under the influence of glacial AcOH. III merely treated in the cold with HCl in absolute alc. gave the blood-red di-HCl salt of 5-p-tolyl-2-oxo-3-methyliminopyrroline (VII). The previously assumed intermediate imido ester, RC(OH):CHC(:NMe)C(:NH)OEt (VIII), corresponding to the nitrile, was isolated as its white HCl salt, which readily changes, even in the absence of air, into the dark red derivative of VII. The distribution of the double bonds shown in III probably occurs only under the influence of the HCl, the free nitrile having the tautomeric structure RCOCH:C(NHMe)CN. The outstanding property of the dark red VII.2HCl is the ease with which the NMe group is replaced by O to form the brick-red 5-p-tolyl-2,3-dioxopyrroline (IX). As with isatin, NaOH cleaves the ring in IX to give α -oxo- γ -imino- γ -ptolylbutyric acid (X) through an intermediate intensely blue alkali salt. Attempts to liberate VII from its HCl salt were unsuccessful. Dilute aqueous alkali or NaHCO3 gave, instead, the yellow-green pseudo base (XI), while excess of concentrated KOH yielded a dark red K salt, C12H11ON2K.2H2O, which regenerated XI with water. NH3 in alc. replaces both the NMe group and the carbonyl O by NH and at the same time 1 mol. alc. is taken up with formation of a product, RC:CH.C(NH2)(OEt).C(:NH).NH (XII), similar in structure to XI; the dark red color immediately produced by HCl shows the ring has not been cleaved. PhNH2 in alc. yields brick-red needles of the 3-phenylimino analog (XIII) of VII. With KOH and also with HCl, XIII forms salts which are red-violet in solution and almost black in the solid state. The HCl salt quant. splits off the HCl at high temps. in vacuo without changing to the brick-red.of the free XIII, showing that the salt formation is accompanied by a simultaneous intramol. rearrangement. In water the HCl salt, like that of VII, is hydrolyzed to IX, but attempts to prepare the pseudo base were unsuccessful; instead was obtained XIII into which the K salt also changes on mere exposure to moist air. This difference in behavior and the very different colors show that the salts of VII and XIII have different structures. As with water and PhNH2, the transfer dark red sait of VII also reacts with compds. having a reactive methylene group. Especially smooth, and under the mildest conditions, is the reaction with CH2(CN)2 to give 5-p-tolyl-2-oxo-3-dicyanomethylenepyrroline (XIV), also obtained from IX or XIII. Surprisingly, XIV forms beautiful violet-black needles and dissolves, although difficultly, in alc. with red-violet color, whereas the corresponding isatin derivative is yellow-red, indicating a fundamental difference in structure. When the alc. solution of XIV is treated with a strong base, it immediately turns steel-blue, but, as with the salts of IX, the blue color quickly disappears and the ring is opened; acids precipitate the yellow cleavage product, RC(NH2):CHC[:C(CN)2]CO2H,

m. 276°, probably in the form of the inner salt, which with boiling

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alc. HCl changes through the intermediate RC(NH2):CHC(CO2H):C(CO2H)C(:NH)O
Et into the compound RC(NH2):CHC(CO2H):CHC(:NH)OEt (XV). X gently heated
with dilute acids yields the dioxo acid. The ring in IX is also cleaved by
piperidine, MeNH2 and NH3 to form compds. of the type RC(:NH)CH2COCONC5H11
(XVI). PhNH2 and CH2(CN)2, on the other hand, react with the 3-CO group,
leaving the ring intact. The similarity of the dioxopyrrolines to isatin
is also shown in their catalytic hydrogenation. There is first formed a
light gray, alc.-insol. product (XVII) corresponding to isatyde which in
the air rapidly regenerates the original compound If the hydrogenation is
continued, the XVII redissolves, and cautious addition of water to the
colorless alc. solution ppts. a completely air-stable crystalline product,
RC(NH2):CHCH(OH)CO2H (XVIII). Reduction of XIII in alc. proceeds 1 step
further, with addition of 2 mols. H and 1 mol. alc. to give the compound
RCH(NH2)CH2CH(NHPh)CO2Et (XIX). For the bearing of the above facts, especially
the color phenomena, on the structures of the mononuclear isatins and
their derivs., the original should be consulted.
Hydroxymethylenepinacolone dioxime (54% yield), m. 84°.
α-tert-Butylisoxazole, b760 156°; its methosulfate with KCN
in water at 0° gave 82% trimethylacetopyruvonitrile methylimide, m.
42°, hydrolyzed by cold concentrated HCl to the pyruvic acid, crystals
with 1 H2O, m. 64°, and by dilute HCl to the amide, m. 115°;
in cold absolute alc. with HCl gas the nitrile imide gave a dark red oil which
with 2 N NaOH or 50% AcOH yielded α-oxo-α-imino-
\delta, \delta-dimethylcaproic acid, m. 185° (gas evolution).
When the red oil was carefully freed from adhering HCl, simple solution in
ordinary alc. resulted in ring cleavage (probably by the water in the
alc.), but AcOEt precipitated a crystalline substance, m. 186°, insol. in all
solvents except alc. and water, which on gentle warming with water gave
trimethylacetopyruvic acid methylimide, m. 183°.
Hydroxymethylenemethyl hexyl ketone oxime, m. 118°.
\alpha-Hexylisoxazole, b11 97-8°, was analyzed as the
chloroplatinate, C20H36O2N2PtCl6, obtained from the methosulfate with
PtCl4. \alpha, \alpha-Dioxodecanonitrile \alpha-methylimide, oil
decomposing on distillation, even in a high vacuum; \alpha, \alpha-
dioxodecanamide, m. 99°. Hydroxymethylene-p-methylacetophenone
oxime (67% yield), m. 133°. II, m. 60°. III, light yellow,
m. 126°. IV (3 g. from 2 g. II and 2.2 mol. MeMgI boiled 2 h. in
ether), yellow, m. 175° (decomposition); heated a short time or allowed
to stand 1 day at room temperature in glacial AcOH, it changed into VI, rhombic
tables, red-brown in incident light, m. 183° (decomposition). V, m.
197°, soluble in AcOH with wine-red color, easily soluble in dilute HCl and
repptd. by NaOH. VII.2HCl (78%), sinters and carbonizes at 183°;
picrate, intensely red, m. 192°. VIII.HCl, from III in cold
dioxane with 0.662 N HCl in absolute alc., decomps. 145°. If in the
treatment of III with alc.-HCl water is present even only in traces the
reaction proceeds in part in an entirely different way, giving in addition to
the dark red salt Me p-toluylpyruvate, m. 84°; free acid, crystals
with 1 H2O, m. 143°. Ag salt of VII, red needles with 1 MeOH,
decomposing 172°. Cu salt, (C12H11ON2)2Cu.4H2O, green, m. 191° (decomposition). XII, m. 153° IX, precipitated quant. in about 6 h. from
VII.2HCl in 20 parts cold water, m. 229-30°; a cold alc. suspension
treated with somewhat less than 1 mol. EtOK-solution at once becomes
blue-violet and soon deposits the K salt, C11H8O2NK.2H2O, which is not
very stable even when dry; one sample had become yellowish after 14 days.
Alc. IX treated with aqueous NaOH also immediately turns blue-violet but the
color rapidly disappears and on cautious acidification X, m. 155°,
seps. Piperidide (XVI), m. 184°. Amide, C11H12O2N2.0.5H2O, m.
179°. Methylamide (0.5H2O), m. 169°. XVIII, turns brown and carbonizes 245-50°. XIII, m. 237°. XIX, m.
123°. XIV was obtained in 92% yield; its melting or decomposition point
is so extraordinarily high that it could not be determined XV.2HCl,
lemon-yellow, m. 148-9°.
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